

**UNITED STATES
SECURITIES AND EXCHANGE COMMISSION**

Washington, D.C. 20549

Form 10-K

(Mark One)

☒ **ANNUAL REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934**

For the fiscal year ended December 31, 2025

or

☐ **TRANSITION REPORT PURSUANT TO SECTION 13 OR 15(d) OF THE SECURITIES EXCHANGE ACT OF 1934**

Commission file number 001-37702

Amgen Inc.

(Exact name of registrant as specified in its charter)

Delaware

(State or other jurisdiction of
incorporation or organization)

**One Amgen Center Drive
Thousand Oaks**

California

(Address of principal executive offices)

95-3540776

(I.R.S. Employer
Identification No.)

91320-1799

(Zip Code)

(805) 447-1000

(Registrant's telephone number, including area code)

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol (s)	Name of each exchange on which registered
Common stock, \$0.0001 par value	AMGN	The Nasdaq Global Select Market
2.00% Senior Notes due 2026	AMGN26	The Nasdaq Global Select Market

Securities registered pursuant to Section 12(g) of the Act: **None**

Indicate by check mark if the registrant is a well-known seasoned issuer, as defined in Rule 405 of the Securities Act. Yes ☒ No ☐

Indicate by check mark if the registrant is not required to file reports pursuant to Section 13 or Section 15(d) of the Act. Yes ☐ No ☒

Indicate by check mark whether the registrant (1) has filed all reports required to be filed by Section 13 or Section 15(d) of the Securities Exchange Act of 1934 during the preceding 12 months (or for such shorter period that the registrant was required to file such reports), and (2) has been subject to such filing requirements for the past 90 days. Yes ☒ No ☐

Indicate by check mark whether the registrant has submitted electronically every Interactive Data File required to be submitted pursuant to Rule 405 of Regulation S-T (§ 232.405 of this chapter) during the preceding 12 months (or for such shorter period that the registrant was required to submit such files). Yes ☒ No ☐

Indicate by check mark whether the registrant is a large accelerated filer, an accelerated filer, a non-accelerated filer, a smaller reporting company, or an emerging growth company. See the definitions of "large accelerated filer," "accelerated filer," "smaller reporting company," and "emerging growth company" in Rule 12b-2 of the Exchange Act.

Large accelerated filer	Accelerated filer	Non-accelerated filer	Smaller reporting company	Emerging growth company
☒	☐	☐	☐	☐

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. ☐

Indicate by check mark whether the registrant has filed a report on and attestation to its management's assessment of the effectiveness of its internal control over financial reporting under Section 404(b) of the Sarbanes-Oxley Act (15 U.S.C. 7262(b)) by the registered public accounting firm that prepared or issued its audit report. ☒

If securities are registered pursuant to Section 12(b) of the Act, indicate by check mark whether the financial statements of the registrant included in the filing reflect the correction of an error to previously issued financial statements. ☐

Indicate by check mark whether any of those error corrections are restatements that required a recovery analysis of incentive-based compensation received by any of the registrant's executive officers during the relevant recovery period pursuant to §240.10D-1(b). ☐

Indicate by check mark whether the registrant is a shell company (as defined in Rule 12b-2 of the Act) Yes ☐ No ☒

The approximate aggregate market value of voting and non-voting stock held by non-affiliates of the registrant was \$150.1 billion as of the last business day of the registrant's most recently completed second fiscal quarter.^(A)

(A) Excludes 879,180 shares of common stock held by directors and executive officers, and any stockholders whose ownership exceeds ten percent of the shares outstanding, as of the last business day of the registrant's most recently completed second fiscal quarter. Exclusion of shares held by any person should not be construed to indicate that such person possesses the power, directly or indirectly, to direct or cause the direction of the management or policies of the registrant, or that such person is controlled by or under common control with the registrant.

539,067,675

(Number of shares of common stock outstanding as of February 10, 2026)

DOCUMENTS INCORPORATED BY REFERENCE

Specified portions of the registrant's Proxy Statement with respect to the 2026 Annual Meeting of Stockholders to be held on May 19, 2026, are incorporated by reference into Part III of this annual report.

INDEX

	<u>Page No.</u>
<u>DEFINED TERMS AND PRODUCTS</u>	<u>ii</u>
<u>PART I</u>	<u>1</u>
Item 1. <u>BUSINESS</u>	<u>1</u>
<u>Significant Developments</u>	<u>1</u>
<u>Marketing, Distribution and Selected Marketed Products</u>	<u>3</u>
<u>Reimbursement</u>	<u>11</u>
<u>Manufacturing, Distribution and Raw Materials</u>	<u>13</u>
<u>Government Regulation</u>	<u>15</u>
<u>Research and Development and Selected Product Candidates</u>	<u>18</u>
<u>Business Relationships</u>	<u>24</u>
<u>Human Capital Resources</u>	<u>25</u>
<u>Information about our Executive Officers</u>	<u>27</u>
<u>Geographic Area Financial Information</u>	<u>28</u>
<u>Investor Information</u>	<u>28</u>
Item 1A. <u>RISK FACTORS</u>	<u>29</u>
Item 1B. <u>UNRESOLVED STAFF COMMENTS</u>	<u>55</u>
Item 1C. <u>CYBERSECURITY</u>	<u>55</u>
Item 2. <u>PROPERTIES</u>	<u>57</u>
Item 3. <u>LEGAL PROCEEDINGS</u>	<u>57</u>
Item 4. <u>MINE SAFETY DISCLOSURES</u>	<u>58</u>
<u>PART II</u>	<u>59</u>
Item 5. <u>MARKET FOR REGISTRANT’S COMMON EQUITY, RELATED STOCKHOLDER MATTERS AND ISSUER PURCHASES OF EQUITY SECURITIES</u>	<u>59</u>
Item 6. <u>RESERVED</u>	<u>60</u>
Item 7. <u>MANAGEMENT’S DISCUSSION AND ANALYSIS OF FINANCIAL CONDITION AND RESULTS OF OPERATIONS</u>	<u>61</u>
Item 7A. <u>QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK</u>	<u>81</u>
Item 8. <u>FINANCIAL STATEMENTS AND SUPPLEMENTARY DATA</u>	<u>83</u>
Item 9. <u>CHANGES IN AND DISAGREEMENTS WITH ACCOUNTANTS ON ACCOUNTING AND FINANCIAL DISCLOSURE</u>	<u>83</u>
Item 9A. <u>CONTROLS AND PROCEDURES</u>	<u>83</u>
Item 9B. <u>OTHER INFORMATION</u>	<u>85</u>
Item 9C. <u>DISCLOSURE REGARDING FOREIGN JURISDICTIONS THAT PREVENT INSPECTIONS</u>	<u>85</u>
<u>PART III</u>	<u>85</u>
Item 10. <u>DIRECTORS, EXECUTIVE OFFICERS AND CORPORATE GOVERNANCE</u>	<u>85</u>
Item 11. <u>EXECUTIVE COMPENSATION</u>	<u>85</u>
Item 12. <u>SECURITY OWNERSHIP OF CERTAIN BENEFICIAL OWNERS AND MANAGEMENT AND RELATED STOCKHOLDER MATTERS</u>	<u>86</u>
Item 13. <u>CERTAIN RELATIONSHIPS AND RELATED TRANSACTIONS AND DIRECTOR INDEPENDENCE</u>	<u>87</u>
Item 14. <u>PRINCIPAL ACCOUNTANT FEES AND SERVICES</u>	<u>87</u>
<u>PART IV</u>	<u>88</u>
Item 15. <u>EXHIBITS AND FINANCIAL STATEMENT SCHEDULES</u>	<u>88</u>
Item 16. <u>FORM 10-K SUMMARY</u>	<u>94</u>
<u>SIGNATURES</u>	<u>95</u>

Defined Terms and Products

Defined terms

We use several terms in this Annual Report on Form 10-K, including but not limited to those that are finance, regulation and disease-state related as well as names of other companies, which are provided below.

Term	Description
2017 Tax Act	Tax Cuts and Jobs Act of 2017
340B Program	Federal 340B Drug Pricing Program
AI	artificial intelligence
Amended 2009 Plan	Amended and Restated 2009 Equity Incentive Plan
AOCI	accumulated other comprehensive income (loss)
AstraZeneca	AstraZeneca plc
ASU	Accounting Standards Update
ATMOS	Amgen Technology and Medical Organizations
B-ALL	B-cell acute lymphoblastic leukemia
BeOne	BeOne Medicines Ltd. (formerly BeiGene, Ltd.)
BiTE®	bispecific T-cell engager
BLA	Biologics License Application
BPCIA	Biologics Price Competition and Innovation Act of 2009
CCPA	California Consumer Privacy Act of 2018
CDT	Cybersecurity & Digital Trust
CGRP	calcitonin gene-related peptide
ChemoCentryx	ChemoCentryx, Inc.
chemotherapy	anticancer medicines
CHMP	Committee for Medicinal Products for Human Use
CIO	Chief Information Officer
CISO	Chief Information Security Officer
CMS	Centers for Medicare & Medicaid Services
COSO	Committee of Sponsoring Organizations of the Treadway Commission
CRCC	Corporate Responsibility and Compliance Committee
DLL3	delta-like ligand 3
DOJ	U.S. Department of Justice
EC	European Commission
EMA	European Medicines Agency
EO	Executive Order
EPO	European Patent Office
EPS	earnings per share
EU	European Union
FASB	Financial Accounting Standards Board
FCPA	U.S. Foreign Corrupt Practices Act
FDA	U.S. Food and Drug Administration
FDCA	Federal Food, Drug, and Cosmetic Act
Fitch	Fitch Ratings, Inc.
GAAP	U.S. generally accepted accounting principles
GDPR	General Data Protection Regulation
GIPR	glucose-dependent insulintropic polypeptide receptor
GLP-1	glucagon like peptide 1
HHS	U.S. Department of Health and Human Services

Term	Description
Horizon	Horizon Therapeutics plc
IGF-1R	insulin-like growth factor-1 receptor
IgG4-RD	Immunoglobulin G4-related disease
IND	Investigational New Drug Application
IPR&D	in-process research and development
IRA	Inflation Reduction Act of 2022
IRS	Internal Revenue Service
July MFN Letter	Letter dated July 31, 2025, by the Administration to a number of pharmaceutical manufacturers, including Amgen
KRAS	Kirsten rat sarcoma viral oncogene homologue
Kyowa Kirin	Kyowa Kirin Co., Ltd.
LDL-C	low-density lipoprotein cholesterol
MAA	Marketing Authorisation Application
MD&A	management's discussion and analysis
MFN	Most-Favored-Nations
MFN EO	Most-Favored-Nations Prescription Drug Pricing Executive Order, dated May 12, 2025
Moody's	Moody's Investors Service, Inc.
MRD	minimal residual disease
Neumora	Neumora Therapeutics, Inc.
NIH	National Institutes of Health
NOL	net operating loss
NSCLC	non-small cell lung cancer
OB3	P.L. 119-21, also The One Big Beautiful Bill Act signed into law on July 4, 2025
OECD	Organisation for Economic Co-operation and Development
PBM	pharmacy benefit manager
PCSK9	proprotein convertase subtilisin/kexin type 9
PDAB	Prescription Drug Affordability Board
R&D	research and development
RANKL	receptor activator of nuclear factor kappa-B ligand
RAR	Revenue Agent Report
RAS	Rat sarcoma viral oncogene
REMS	Risk Evaluation and Mitigation Strategy
ROU	right-of-use
ROW	rest of world
RSUs	restricted stock units
S&P	Standard & Poor's Financial Services LLC
SEC	U.S. Securities and Exchange Commission
SG&A	selling, general and administrative
SOFR	Secured Overnight Financing Rate
TED	thyroid eye disease
Teneobio	Teneobio, Inc.
U.S. Treasury	U.S. Department of the Treasury
USPTO	U.S. Patent and Trademark Office
UTB	unrecognized tax benefit

Products

The brand names of our products, our delivery devices and certain of our product candidates and their associated generic names are provided below.

Term	Description
ACTIMMUNE	ACTIMMUNE® (interferon gamma-1b)
Aimovig	Aimovig® (erenumab-aooe)
AMJEVITA/AMGEVITA	AMJEVITA® (adalimumab-atto)/AMGEVITA™ (adalimumab)
Aranesp	Aranesp® (darbepoetin alfa)
AutoTouch	AutoTouch®
AVSOLA	AVSOLA® (infliximab-axxq)
BKEMV/BEKEMV	BKEMV® (eculizumab-aeeb)/BEKEMV™ (eculizumab)
BLINCYTO	BLINCYTO® (blinatumomab)
BUPHENYL	BUPHENYL® (sodium phenylbutyrate)
ConfiPen	ConfiPen™
Corlanor	Corlanor® (ivabradine)
ENBREL	Enbrel® (etanercept)
ENBREL Mini	ENBREL Mini®
EPOGEN	EPOGEN® (epoetin alfa)
EVENITY	EVENITY® (romosozumab-aqqg)
IMDELLTRA/IMDYLLTRA	IMDELLTRA® (tarlatamab-dlle)/IMDYLLTRA™ (tarlatamab)
IMLYGIC	IMLYGIC® (talimogene laherparepvec)
KANJINTI	KANJINTI® (trastuzumab-anns)
KRYSTEXXA	KRYSTEXXA® (pegloticase)
KYPROLIS	KYPROLIS® (carfilzomib)
LUMAKRAS/LUMYKRAS	LUMAKRAS®/LUMYKRAS™ (sotorasib)
MariTide	Maridebart cafraglutide (MariTide™)
MVASI	MVASI® (bevacizumab-awwb)
Neulasta	Neulasta® (pegfilgrastim)
NEUPOGEN	NEUPOGEN® (filgrastim)
Nplate	Nplate® (romiplostim)
Onpro	Onpro®
Otezla	Otezla® (apremilast)
Parsabiv	Parsabiv® (etelcalcetide)
PAVBLU	PAVBLU® (afibercept-ayyh)
PENNSAID	PENNSAID® (diclofenac sodium topical solution) 2%
PROCYSBI	PROCYSBI® (cysteamine bitartrate)
Prolia	Prolia® (denosumab)
QUINSAIR	QUINSAIR® (levofloxacin)
RAVICTI	RAVICTI® (glycerol phenylbutyrate)
RAYOS	RAYOS® (prednisone)
Repatha	Repatha® (evolocumab)
RIABNI	RIABNI® (rituximab-arrx)
Sensipar/Mimpara	Sensipar®/Mimpara™ (cinacalcet)
SureClick	SureClick®
TAVNEOS	TAVNEOS® (avacopan)
TEPEZZA	TEPEZZA® (teprotumumab-trbw)

Term	Description
TEZSPIRE	TEZSPIRE [®] (tezepelumab-ekko)
UPLIZNA	UPLIZNA [®] (inebilizumab-cdon)
Vectibix	Vectibix [®] (panitumumab)
WEZLANA/WEZENLA	WEZLANA [®] (ustekinumab-auub)/WEZENLA [™] (ustekinumab)
XGEVA	XGEVA [®] (denosumab)

Products referenced in this report that are not included in the above list are the properties of their respective owners.

PART I

Item 1. BUSINESS

Amgen Inc. (including its subsidiaries, referred to as “Amgen,” “the Company,” “we,” “our” or “us”) discovers, develops, manufactures and delivers innovative medicines to fight some of the world’s toughest diseases. We focus on areas of high unmet medical need and leverage our expertise to strive for solutions that dramatically improve people’s lives, while also reducing the social and economic burden of disease. We helped launch the biotechnology industry more than 45 years ago and have grown to be one of the world’s leading independent biotechnology companies. Our robust pipeline includes potential first-in-class medicines at all stages of development. We have a presence in approximately 100 countries worldwide.

Amgen was incorporated in California in 1980 and became a Delaware corporation in 1987. Amgen operates in one operating segment: human therapeutics.

Significant Developments

Following is a summary of significant developments affecting our business that have occurred and that we have reported since the filing of our Annual Report on Form 10-K for the year ended December 31, 2024.

Products/Pipeline

Repatha

In August 2025, we announced that the FDA broadened the approved use of Repatha to include adults at increased risk for major adverse cardiovascular events (MACE) due to uncontrolled low-density lipoprotein cholesterol (LDL-C), removing the previous requirement that a patient have been diagnosed with cardiovascular (CV) disease.

In November 2025, we announced detailed results from the Phase 3 VESALIUS-CV clinical trial, which showed that Repatha achieved statistically significant and clinically meaningful reductions in MACEs in high-risk adults without a prior heart attack or stroke, when added to statins or other LDL-C-lowering treatments. Repatha demonstrated a 25% relative reduction in the risk of a composite of coronary heart disease (CHD) death, heart attack or ischemic stroke (3-P MACE), and a 19% reduction in a broader composite that also included any ischemia-driven arterial revascularization (4-P MACE). Repatha also reduced the risk of heart attack by 36%. In a cohort of patients included in a lipid sub-study, the median achieved LDL-C was 45 mg/dL compared to 109 mg/dL in the placebo arm. No new safety signals were identified, and tolerability was consistent with the current prescribing information in the United States.

UPLIZNA

In April 2025, we announced the FDA approved UPLIZNA for the treatment of Immunoglobulin G4-related disease (IgG4-RD) in adult patients. UPLIZNA is the first and only FDA approved treatment for adults living with IgG4-RD.

In December 2025, we announced the FDA approved UPLIZNA for the treatment of generalized myasthenia gravis (gMG) in adults who are anti-acetylcholine receptor (AChR) and anti-muscle specific tyrosine kinase (MuSK) antibody positive.

IMDELLTRA/IMDYLLTRA

In June 2025, we announced interim results from the global Phase 3 DeLLphi-304 trial evaluating IMDELLTRA/IMDYLLTRA in patients with small cell lung cancer (SCLC) who had progressed on or after one line of platinum-based chemotherapy. The study demonstrated that IMDELLTRA/IMDYLLTRA significantly reduced the risk of death by 40% compared to standard-of-care chemotherapy, with a median overall survival of 13.6 months compared to 8.3 months. Additionally, IMDELLTRA/IMDYLLTRA showed a statistically significant improvement in median progression-free survival of 4.2 months compared to 3.7 months and enhanced patient-reported outcomes related to cancer-associated symptoms, including dyspnea and cough. The safety profile of IMDELLTRA/IMDYLLTRA was consistent with prior studies.

In November 2025, we announced that the FDA granted full approval for IMDELLTRA for the treatment of adult patients with extensive stage small cell lung cancer (ES-SCLC) with disease progression on or after platinum-based chemotherapy, converting its prior accelerated approval to full approval based on results from the global Phase 3 DeLLphi-304 study.

TEZSPIRE

In October 2025, we announced that the FDA approved TEZSPIRE as an add-on maintenance treatment of inadequately controlled chronic rhinosinusitis with nasal polyps (CRSwNP) in adult and pediatric patients aged 12 years and older.

TEPEZZA

In June 2025, the European Commission granted marketing authorization approval of TEPEZZA for treatment of adults with moderate-to-severe thyroid eye disease (TED).

Maridebart cafraglutide (MariTide™)

In March 2025, we announced the initiation of two global Phase 3 studies to evaluate MariTide, a differentiated antibody-peptide conjugate that activates the glucagon like peptide 1 (GLP-1) receptor and antagonizes the glucose-dependent insulinotropic polypeptide receptor (GIPR), in chronic weight management: one study in adults living with obesity or overweight without Type 2 diabetes and another study in adults living with obesity or overweight with Type 2 diabetes.

In June 2025, the underlying details from Part 1 of the Phase 2 study of MariTide and complete results from the primary analysis of the Phase 1 pharmacokinetics low dose initiation (PK-LDI) study evaluating lower starting doses of MariTide were presented at the American Diabetes Association 85th Scientific Sessions and simultaneously published in *The New England Journal of Medicine*.

In November 2025, we announced that, in addition to the two global Phase 3 studies announced in March 2025, four global Phase 3 studies were underway, with studies in adults living with established atherosclerotic cardiovascular disease and obesity or overweight; in adults living with heart failure with preserved or mildly reduced ejection fraction and obesity; in adults living with obstructive sleep apnea on positive airway pressure therapy and living with obesity or overweight; and in adults living with obstructive sleep apnea not on positive airway pressure therapy and living with obesity or overweight.

In January 2026, we announced that Part 2 of the Phase 2 chronic weight management study, an exploratory evaluation of MariTide treatment for an additional 52 weeks in people who lost at least 15% of their body weight in the 52-week Part 1 of the Phase 2 chronic weight management study, is complete. Key findings included the following: the large majority of participants maintained the weight loss achieved in Part 1 for an additional 52 weeks on a lower monthly dose or quarterly dose of MariTide; the second year of MariTide treatment was very well tolerated, including at quarterly doses, with a very low incidence of nausea and vomiting and no new safety signals observed; and improvements in cardiometabolic parameters were sustained with MariTide at effective maintenance doses for a full second year.

In January 2026 we also announced that a Phase 2 study of MariTide for the treatment of Type 2 diabetes in adults living with and without obesity has completed the 24-week timepoint. Key findings included the following: robust and clinically meaningful reduction in both hemoglobin A1c (HbA1c) and weight with monthly MariTide at 24 weeks, which is in line with results seen in the Type 2 diabetes population in Part 1 of the Phase 2 chronic weight management study, at 24 weeks; safety and tolerability profile consistent with the GLP-1 class, where the most common side effects were gastrointestinal-related, predominantly mild-to-moderate in nature, and occurred primarily during dose escalation; and favorable improvement in cardiometabolic parameters.

TAVNEOS

TAVNEOS was approved by the FDA in October 2021 for the adjunctive treatment of adult patients with severe active anti-neutrophil cytoplasmic autoantibody (ANCA)-associated vasculitis (AAV) in combination with standard therapy including glucocorticoids. TAVNEOS was developed by ChemoCentryx. Amgen acquired ChemoCentryx in October 2022, after TAVNEOS had been on the market for a year. On January 16, 2026, the FDA requested that ChemoCentryx voluntarily withdraw TAVNEOS from the U.S. market. The FDA raised concerns about the process followed by ChemoCentryx to re-adjudicate primary endpoint results for 9 of the 331 patients in its pivotal clinical trial. Hepatotoxicity, which is a known infrequent risk of TAVNEOS treatment for AAV, was also raised in the context of the benefit-risk profile of the medicine. Amgen is not aware of any issue with the underlying patient data from the ChemoCentryx clinical trial, and after review of the relevant clinical data and years of real-world evidence, Amgen is confident that TAVNEOS demonstrates effectiveness and a favorable benefit-risk profile. On January 28, 2026, following FDA regulatory process, Amgen informed the FDA that it did not intend to withdraw TAVNEOS from the market. Amgen is evaluating next steps with the FDA to determine a path forward, while keeping patient safety, needs and support at the forefront.

Bemarituzumab

In February 2026, we announced that, based upon data from the FORTITUDE-101 and FORTITUDE-102 Phase 3 studies, the Company does not intend to pursue regulatory approval of bemarituzumab in first-line gastric cancer.

Rocatinlimab

In January 2026, we and Kyowa Kirin agreed to terminate the rocatinlimab collaboration agreement and to transition control of the global development and commercialization program to Kyowa Kirin, subject to receipt of clearance under the

Hart-Scott-Rodino Antitrust Improvements Act of 1976. In February 2026, we received such regulatory clearance. In the first quarter of 2026, Kyowa Kirin will assume full responsibility for rocatinlimab worldwide, except that Amgen will continue to manufacture rocatinlimab and perform other transition activities for an agreed upon period of time.

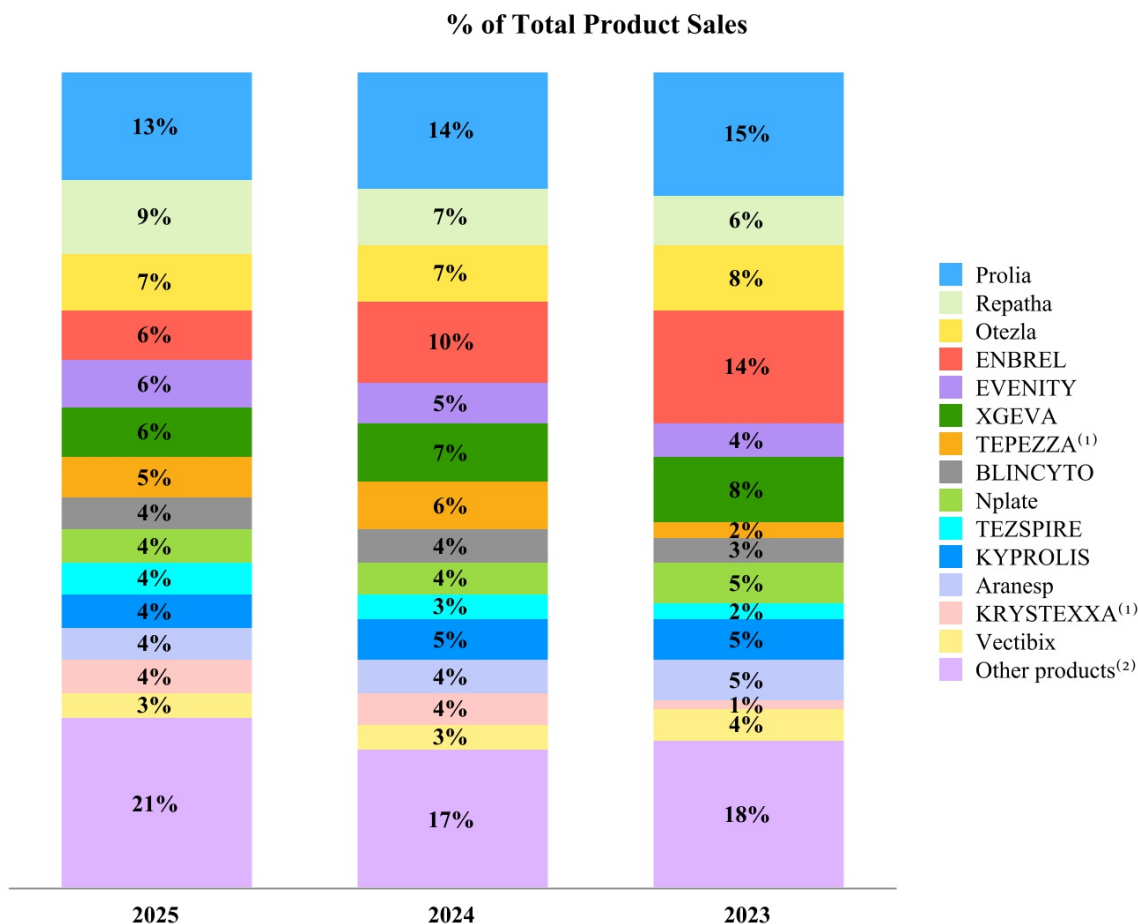
Marketing, Distribution and Selected Marketed Products

The largest concentration of our sales and marketing forces is based in the United States and Europe. We also commercialize and market our products into other geographic territories, including Japan, China and other parts of Asia, Latin America and the Middle East by using our own affiliates, by acquiring existing third-party businesses or product rights or by collaborating with third parties. In the Asia Pacific region, we also sell our products in partnership with other companies, including Astellas Pharma Inc., BeOne, Mitsubishi Tanabe Pharma Corporation, Takeda Pharmaceutical Co., Ltd. and Kyowa Kirin. This international footprint allows us to deliver our medicines to more patients globally. See Business Relationships for our significant alliances. Whether we use our own sales and marketing forces or a third party's services varies across these markets. Such use typically depends on several factors, including the nature of entry into the new market, the size of an opportunity and operational capabilities. Together with our collaborators, we market our products to healthcare providers, including physicians or their clinics, dialysis centers, hospitals and pharmacies.

In the United States, substantially all of our sales are to pharmaceutical wholesale distributors, which is the principal means of distributing our products to healthcare providers. We market certain products through direct-to-patient channels, including print, television and online media. For further discussion, see Government Regulation—*Regulation in the United States—Regulation of Product Marketing and Promotion*. Outside the United States, we sell principally to healthcare providers and/or pharmaceutical wholesale distributors depending on the distribution practice in each country.

Our product sales to three large wholesalers, McKesson Corporation, Cencora, Inc. and Cardinal Health, Inc., each individually accounted for more than 10% of total revenues for each of the years 2025, 2024 and 2023. On a combined basis, these wholesalers accounted for 77%, 77% and 79% of worldwide gross revenues for 2025, 2024 and 2023, respectively. We monitor the financial condition of our larger customers and limit our credit exposure by setting credit limits and, in certain circumstances, by requiring letters of credit or obtaining credit insurance.

Our products are marketed around the world, with the United States as our largest market. The following chart shows our product sales by principal product, and the table below (dollar amounts in millions) shows product sales by geography for the years 2025, 2024 and 2023.



	2025		2024		2023	
Product Sales by Geography:						
U.S.	\$	25,656	73 %	\$	23,301	72 %
ROW		9,492	27 %		8,725	27 %
Total	\$	35,148	100 %	\$	32,026	100 %

⁽¹⁾ TEPEZZA and KRYSTEXXA were acquired from our Horizon acquisition on October 6, 2023, and include product sales in the periods after the acquisition date.

⁽²⁾ Consists of product sales of our non-principal products.

Prolia

We market Prolia in many countries around the world. Prolia and XGEVA contain the same active ingredient but are approved for different indications, patient populations, dose and frequency of administration. Prolia was launched in the United States and Europe in 2010. In the United States, it is used primarily in the indication for the treatment of postmenopausal women with osteoporosis at high risk of fracture and for treatment to increase bone mass in men with osteoporosis at high risk of fracture. In Europe, Prolia is used primarily for the treatment of osteoporosis in men and postmenopausal women at increased risk of fracture. Our patents for RANKL antibodies, including sequences, for Prolia expired in February 2025 in the United States and in November 2025 in select countries in Europe.

Repatha

We market Repatha, a PCSK9 inhibitor, in many countries around the world. Repatha was launched in 2015 and is indicated to reduce the risk of major adverse cardiovascular (CV) events (CV death, myocardial infarction, stroke, unstable angina requiring hospitalization, or coronary revascularization) in adults at increased risk for these events. Repatha is also indicated to reduce LDL-C in adults with primary hyperlipidemia, including heterozygous familial hypercholesterolemia (HeFH). See Significant Developments for additional information regarding regulatory developments.

Otezla

We market Otezla, a small molecule that inhibits phosphodiesterase 4 (PDE4), in many countries around the world. Otezla was acquired from Bristol Myers Squibb Company in November 2019 after its acquisition of Celgene Corporation. Otezla is an oral therapy approved for the treatment of adults with plaque psoriasis across all severities (in the United States, Japan and Australia) and moderate-to-severe plaque psoriasis (in other global markets, including Europe); for adults with active psoriatic arthritis; for adults with oral ulcers associated with Behçet's disease; and for pediatric patients six years of age and older and weighing at least 20 kilograms with moderate-to-severe plaque psoriasis who are candidates for phototherapy or systemic therapy. Otezla is also approved in Japan for the treatment of palmoplantar pustulosis (PPP).

ENBREL

We market ENBREL, a tumor necrosis factor blocker, in the United States and Canada. ENBREL was launched in 1998 and is used primarily in indications for the treatment of adult patients with moderately-to-severely active rheumatoid arthritis, patients with chronic moderate-to-severe plaque psoriasis who are candidates for systemic therapy or phototherapy and patients with active psoriatic arthritis.

EVENTY

Together with our collaboration partners, we market EVENTY in many countries around the world. EVENTY was launched in the United States and Japan in 2019. In the United States, it is used in the indication for the treatment of osteoporosis in postmenopausal women at high risk for fracture, defined as a history of osteoporotic fracture, or multiple risk factors for fracture; or patients who have failed or are intolerant to other available osteoporosis therapy. In Japan, EVENTY is used primarily in the indication for the treatment of osteoporosis in men and postmenopausal women at high risk of fracture.

XGEVA

We market XGEVA in many countries around the world. Prolia and XGEVA contain the same active ingredient but are approved for different indications, patient populations, dose and frequency of administration. XGEVA was launched in 2010 and is used primarily in the indication for prevention of skeletal-related events (pathological fracture, radiation to bone, spinal cord compression or surgery to bone) in patients with bone metastases from solid tumors and multiple myeloma. Our patents for RANKL antibodies, including sequences, for XGEVA expired in February 2025 in the United States and in November 2025 in select countries in Europe.

TEPEZZA

We market TEPEZZA for the treatment of TED in multiple countries globally, including the United States, Japan, Brazil, among others. TEPEZZA was acquired through our Horizon acquisition in October 2023. TEPEZZA is a fully human monoclonal antibody and a targeted inhibitor of the insulin-like growth factor-1 receptor (IGF-1R) that is the first and only approved medicine for the treatment of TED. See Significant Developments for additional information regarding regulatory developments.

BLINCYTO

We market BLINCYTO in many countries around the world. BLINCYTO was launched in 2014 and has proven efficacy in a wide range of patients with CD19-positive B-cell acute lymphoblastic leukemia (B-ALL), including those who are MRD(–) or MRD(+) in frontline consolidation, and those with relapsed or refractory (R/R) disease. Acute lymphoblastic leukemia (ALL) is a cancer of the blood in which a particular kind of white blood cell is growing out of control.

Nplate

We market Nplate in many countries around the world. Nplate was launched in 2008 and is indicated to treat thrombocytopenia in patients with immune thrombocytopenia (ITP) who have had an insufficient response to corticosteroids, immunoglobulins or splenectomy.

TEZSPIRE

Together with our collaboration partner, we market TEZSPIRE in many countries around the world. TEZSPIRE is currently approved for the treatment of severe uncontrolled asthma (SUA) in the United States, Europe, Japan and more than 70 countries across the globe. In addition, TEZSPIRE is approved for chronic rhinosinusitis with nasal polyps (CRSwNP) in the United States, Europe and several additional countries. TEZSPIRE is a first-in-class human monoclonal antibody that works on the primary source of inflammation: the airway epithelium, which is the first point of contact for viruses, allergens, pollutants and other environmental insults. See Significant Developments for additional information regarding regulatory developments.

KYPROLIS

We market KYPROLIS primarily in the United States and Europe. KYPROLIS was launched in 2012 and is indicated in combination with (i) dexamethasone, (ii) lenalidomide plus dexamethasone, (iii) daratumumab plus dexamethasone, (iv) daratumumab plus hyaluronidase-fihj plus dexamethasone, and (v) isatuximab plus dexamethasone for the treatment of patients with relapsed or refractory multiple myeloma who have received one to three prior lines of therapy. It is also approved as a single agent for patients with relapsed or refractory multiple myeloma who have received one or more previous therapies.

Aranesp

We market Aranesp primarily in the United States and Europe. Aranesp was launched in 2001 and is indicated to treat a lower-than-normal number of red blood cells (anemia) caused by chronic kidney disease (CKD) in both patients on dialysis and patients not on dialysis. Aranesp is also indicated for anemia in certain patients with non-myeloid malignancies who are receiving concomitant myelosuppressive chemotherapy.

KRYSTEXXA

We market KRYSTEXXA in the United States. KRYSTEXXA was acquired through our Horizon acquisition in October 2023. KRYSTEXXA is the first and only FDA-approved medicine for the treatment of chronic refractory gout.

Vectibix

We market Vectibix in many countries around the world. Vectibix was launched in 2006 and is indicated for the treatment of patients with wild-type RAS metastatic colorectal cancer (mCRC, which is a cancer that has spread outside the colon and rectum) and in the United States, in combination with LUMAKRAS, for the treatment of adult patients with KRAS G12C-mutated mCRC, who have received prior fluoropyrimidine-, oxaliplatin- and irinotecan-based chemotherapy. RAS status is determined by an FDA-approved test.

Other Marketed Products

We also market a number of other products in various markets worldwide, including but not limited to MVASI, PAVBLU, UPLIZNA, IMDELLTRA/IMDYLLTRA, AMJEVITA/AMGEVITA, TAVNEOS, Neulasta, LUMAKRAS/LUMYKRAS, RAVICTI, Parsabiv, Aimovig, WEZLANA/WEZENLA and PROCYSBI.

Patents

The following table lists our outstanding material patents for the indicated product by territory, general subject matter and latest expiry date. Certain of the European patents are subjects of supplemental protection certificates that provide additional protection for the products in certain European countries beyond the dates listed in the table. See footnotes to the patent table below.

One or more patents with the same or earlier expiry dates may fall under the same general subject matter and are not listed separately.

Product	Territory	General subject matter	Expiration
Repatha® (evolocumab)	U.S.	Antibodies	8/27/2029
	U.S.	Methods of treatment	5/10/2032
	Europe	Compositions ⁽¹⁾	8/22/2028
	Europe	Methods of treatment	5/10/2032
	Europe	Formulation	5/3/2033
Otezla®/Otezla XR™ (apremilast)	U.S.	Compositions and compounds ⁽²⁾	2/16/2028
	U.S.	Oral dosage forms for extended release and methods of treatment	5/27/2042
Enbrel® (etanercept)	U.S.	Fusion protein and pharmaceutical compositions	11/22/2028
	U.S.	DNA encoding fusion protein and methods of making fusion protein	4/24/2029
	U.S.	Formulations and methods of preparing formulations	10/19/2037
	U.S.	Methods of treatment using formulations	2/22/2038
EVENITY® (romosozumab-aqqg)	U.S.	Antibodies	4/25/2026
	U.S.	Formulation and methods of using formulation	5/11/2031
	U.S.	Methods of treatment	4/9/2033
	Europe	Antibodies ⁽¹⁾	4/28/2026
	Europe	Formulation and methods of using formulation	5/11/2031
	Europe	Methods of treatment	4/18/2032
TEPEZZA® (teprotumumab-trbw) ⁽³⁾	U.S.	IGF-1R antibodies	3/3/2029
	U.S.	Methods of treatment	12/11/2039
BLINCYTO® (blinatumomab)	U.S.	Pharmaceutical compositions and bifunctional polypeptides	4/6/2030
	U.S.	Method of treatment	8/26/2031
	Europe	Bifunctional polypeptides ⁽¹⁾	11/26/2024
	Europe	Method of treatment	11/6/2029
Nplate® (romiplostim)	U.S.	Formulation	2/12/2028
	Europe	Formulation	4/20/2027
TEZSPIRE® (tezepelumab-ekko)	U.S.	Polypeptides	2/3/2034
	U.S.	Methods of treatment	8/23/2038
	Europe	Polypeptides ⁽¹⁾	9/9/2028
KYPROLIS® (carfilzomib)	U.S.	Compositions and compounds ⁽⁴⁾	12/7/2027
	U.S.	Methods of making	5/8/2033
	Europe	Compositions, compounds and methods of treatment ⁽¹⁾	12/7/2025
	Europe	Pharmaceutical compositions and methods of making	5/8/2033
KRYSTEXXA® (pegloticase)	U.S.	Polypeptides and pharmaceutical compositions	4/11/2026
	U.S.	Methods of treatment, including combination therapy	10/5/2040
UPLIZNA® (inebilizumab-cdon)	U.S.	CD19 antibodies and pharmaceutical compositions	6/11/2034
	Europe	CD19 antibodies, pharmaceutical compositions and methods of treatment ⁽¹⁾	9/7/2027
IMDELLTRA® (tarlatamab-dlle)/IMDYLLTRA™ (tarlatamab)	U.S.	Bifunctional polypeptides ⁽⁵⁾	8/12/2036
	Europe	Bifunctional polypeptides	8/1/2036
TAVNEOS® (avacopan)	U.S.	Compounds and pharmaceutical compositions	1/6/2034
	U.S.	Formulations	11/27/2039
	U.S.	Amorphous forms and pharmaceutical compositions and uses thereof	11/27/2039
	Europe	Compounds, pharmaceutical compositions, and methods of treatment ⁽¹⁾	12/21/2029
	Europe	Formulations	11/27/2039
LUMAKRAS®/LUMYKRAS™ (sotorasib)	U.S.	Compounds and pharmaceutical compositions	5/21/2038
	U.S.	Crystalline form, pharmaceutical compositions and methods of treatment	5/20/2040
	U.S.	Methods of treatment	9/15/2040
	Europe	Compounds, pharmaceutical compositions and methods of treatment	5/21/2038

Product	Territory	General subject matter	Expiration
Parsabiv® (etelcalcetide)	U.S.	Compound and pharmaceutical composition	2/7/2031
	U.S.	Formulation	6/27/2034
	U.S.	Methods of making	8/9/2035
	Europe	Compound and pharmaceutical composition ⁽¹⁾	7/29/2030
	Europe	Formulation	6/27/2034
	Europe	Methods of making	4/3/2035
Aimovig® (erenumab-aooe)	U.S.	Polynucleotides encoding CGRP receptor antibodies and methods of making antibodies	12/11/2031
	U.S.	CGRP receptor antibodies	5/17/2032
	U.S.	Methods of treatment	4/22/2036
	U.S.	Compositions and pharmaceutical formulations	4/1/2039
	Europe	CGRP receptor antibodies ⁽¹⁾	12/18/2029
	Europe	Methods of treatment	8/10/2035

⁽¹⁾ A European patent with this subject matter may also be entitled to supplemental protection in one or more countries in Europe, and the length of any such extension will vary by country. For example, supplementary protection certificates have been issued related to the indicated products for patents in at least the following countries:

- evolocumab — France, Italy, Spain and the United Kingdom, expiring in 2031
- romosozumab — France, Germany, Italy, Spain and the United Kingdom, expiring in 2031
- blinatumomab — France, Germany, Italy, Spain and the United Kingdom, expiring in 2029
- tezepelumab — France, Italy and Spain, expiring in 2033
- carfilzomib — France, Germany, Italy, Spain and the United Kingdom, expiring in 2030
- inebilizumab — France, Germany, Italy and Spain, expiring in 2032
- avacopan — France, Italy, Spain and the United Kingdom, expiring in 2034
- etelcalcetide — France, Germany, Italy, Spain and the United Kingdom, expiring in 2031
- erenumab — France, Germany, Italy, Spain and the United Kingdom, expiring in 2033

⁽²⁾ Pediatric exclusivity granted to August 16, 2028 for a patent with this subject matter.

⁽³⁾ We have biologic exclusivity in the United States and regulatory exclusivity in Europe covering teprotumumab-trbw that will expire in 2032 and 2035, respectively.

⁽⁴⁾ Pediatric exclusivity granted to June 7, 2028 for a patent with this subject matter.

⁽⁵⁾ A patent with this subject matter may be entitled to patent term extension in the United States.

Competition

We operate in a highly competitive environment. A number of our marketed products are indicated for disease areas in which other products or treatments are currently available or are being pursued by our competitors through R&D activities. Additionally, some competitor-marketed products target the same genetic pathways as our recently launched marketed products or product candidates. This competition could impact the pricing and market share of our products. We continue to pursue ways of increasing the value of our medicines through innovations, which can include expanding the disease areas for which our products are indicated and finding new methods to make the delivery or manufacture of our medicines easier and less costly. Such activities can offer important opportunities for differentiation. We plan to continue pursuing innovation efforts to strengthen our competitive position. Such position may be based on, among other things, safety, efficacy, reliability, availability, patient convenience, delivery devices, price, reimbursement, access to and timing of market entry and patent position and expiration.

Certain of the existing patents on our principal products have expired, and we face new and increasing competition, including from biosimilars and generics. A biosimilar is another version of a biological product for which marketing approval is sought or has been obtained based on a demonstration that it is “highly similar” to the original reference product. We have experienced adverse effects from biosimilar competition on our originator product sales. Companies have launched versions of EPOGEN, NEUPOGEN, Neulasta and ENBREL (Canada only) with U.S. ENBREL biosimilars approved but not launched. Once multiple biosimilar versions of one of our originator products have launched, competition intensifies rapidly, resulting in accelerated net price declines for both the reference and the biosimilar products. For example, our patents for RANKL antibodies, including sequences, for Prolia and XGEVA expired in February 2025 in the United States and in November 2025 in select countries in Europe, and we expect accelerated sales erosion driven by increased competition, as multiple biosimilars have launched in the United States and ROW. See also Government Regulation—*Regulation in the United States—Approval of Biosimilars*.

We also have our own biosimilar products both in the United States and ROW that are competing against branded and biosimilar versions of our competitors’ products. Since 2018, we have launched eight biosimilars, including the 2025 U.S. launches of WEZLANA, a biosimilar to STELARA®, and BKEMV, a biosimilar to SOLIRIS®. We expect additional biosimilar competition against both our branded and biosimilar products in the future across markets. Although biosimilars compete on price, we believe many patients, providers and payers will continue to place high value on the reputation, supply reliability and safety of our products. As additional biosimilar competitors come to market, we will continue to leverage our global experience to distinguish against both branded and biosimilar competitors.

Although most of our products are biologics, some are small molecule products, including Otezla, KYPROLIS and LUMAKRAS/LUMYKRAS. Because the FDA approval process permits generic manufacturers to rely on the safety and efficacy data of the innovator product rather than having to conduct their own costly and time-consuming clinical trials, generic manufacturers can often develop and market their competing versions of our small molecule products at much lower prices. For example, following loss of exclusivity of patents directed to cinacalcet, the active ingredient in our small molecule calcimimetic Sensipar, we lost a significant share of the market and corresponding revenues in a very short period of time.

The introduction of new products, the development of new processes or technologies by competitors or the emergence of new information about existing products may result in (i) increased competition for our marketed products, even for those protected by patents and/or (ii) reductions in the prices we receive from selling our products. In addition, the development of new treatment options or standards of care may reduce the use of our products or may limit the utility and application of ongoing clinical trials of our product candidates. (As used in this document, the term *clinical trials* may include prospective clinical trials, observational studies, registries and other studies.) See Item 1A. Risk Factors—*Our products face substantial competition and our product candidates are also likely to face substantial competition* and Item 1A. Risk Factors—*We currently face competition from biosimilars and generics and expect to face increasing competition from biosimilars and generics in the future*.

The following table reflects our significant competitors for our principal products and is not exhaustive.

Product	Territory	Competitor-marketed product	Competitors
Prolia	U.S., Europe & Asia Pacific	Bisphosphonates, including generics	Various
	U.S., Europe & Asia Pacific	Denosumab biosimilars	Various
Repatha	U.S., Europe & Asia Pacific	LEQVIO®	Novartis AG
	U.S., Europe & Asia Pacific	PRALUENT®	Regeneron Pharmaceuticals, Inc. Sanofi
Otezla	U.S. & Europe	Skyrizi®	AbbVie Inc.
	U.S. & Europe	Taltz®	Eli Lilly and Company
	U.S. & Europe	Tremfya®	Johnson & Johnson Innovative Medicine ⁽¹⁾
	U.S. & Europe	HUMIRA® ⁽²⁾	AbbVie Inc.
	U.S. & Europe	Cosentyx®	Novartis AG
	U.S. & Europe	SOTYKTU®	Bristol Myers Squibb Company
	U.S. & Europe	Topical products	Various
ENBREL	U.S.	HUMIRA® ⁽²⁾	AbbVie Inc.
	U.S.	RINVOQ®	AbbVie Inc.
	U.S.	ORENCIA®	Bristol Myers Squibb Company
	U.S.	Xeljanz®	Pfizer Inc.
	U.S.	Cosentyx®	Novartis AG
	Canada	Etanercept biosimilars	Various
EVENITY	U.S.	Bisphosphonates, including generics	Various
	Japan	Teribone™	Asahi Kasei Pharma
XGEVA	U.S. & Europe	Zoledronate generics	Various
	U.S., Europe & Asia Pacific	Denosumab biosimilars	Various
BLINCYTO	U.S. & Europe	BESPONSA®	Pfizer Inc.
	U.S. & Europe	Chemotherapy regimen	Various
Nplate	U.S. & Europe	PROMACTA®/REVOLADE™	Novartis AG
TEZSPIRE	U.S.	DUPIXENT®	Regeneron Pharmaceuticals, Inc. Sanofi
	U.S.	FASENRA®	AstraZeneca
	U.S.	NUCALA®	GSK plc.
	U.S.	XOLAIR®	Genentech, Inc. Novartis AG
KYPROLIS	U.S. & Europe	DARZALEX®	Johnson & Johnson Innovative Medicine ⁽¹⁾
	U.S. & Europe	POMALYST®/IMNOVID®	Celgene Corporation ⁽³⁾
	U.S. & Europe	REVLIMID® ⁽⁴⁾	Various
	U.S.	VELCADE®	Takeda Oncology ⁽⁵⁾
	U.S.	CARVYKTI®	Johnson & Johnson Innovative Medicine ⁽¹⁾ Legend Biotech
Aranesp	U.S.	PROCRT® ⁽⁶⁾	Johnson & Johnson Innovative Medicine ⁽¹⁾
	U.S. & Europe	Epoetin alfa biosimilars	Various

Vectibix	U.S. & Europe	Avastin [®]	F. Hoffmann-La Roche Ltd. (Roche)
	U.S. & Europe	ERBITUX [®]	Eli Lilly and Company Merck KGaA
	U.S.	KEYTRUDA [®]	Merck & Co., Inc.
	U.S. & Europe	Chemotherapy regimen	Various

⁽¹⁾ A subsidiary of Johnson & Johnson.

⁽²⁾ Approved biosimilars for HUMIRA[®] available.

⁽³⁾ A subsidiary of Bristol-Myers Squibb Company.

⁽⁴⁾ REVLIMID[®] also includes generics.

⁽⁵⁾ A subsidiary of Takeda Pharmaceutical Co., Ltd.

⁽⁶⁾ PROCIT[®] competes with Aranesp in supportive cancer care and predialysis settings.

TEPEZZA and KRYSTEXXA currently do not face any direct competitors in the United States, Europe or Japan. TEPEZZA faces competition from other therapies, such as corticosteroids, which have been used on an off-label basis to alleviate some of the symptoms of TED. TEPEZZA and KRYSTEXXA may face competition from competitor medicines currently in clinical trials. See TEPEZZA and KRYSTEXXA sections above and Government Regulation—*Regulation of Orphan Medicines*.

Reimbursement

Sales of our products are dependent on the availability and extent of coverage and reimbursement from third-party payers. In many markets around the world, these payers, including government health systems, private health insurers and other organizations, remain focused on reducing the cost of healthcare; and their efforts have intensified, in part, as a result of uncertain macroeconomic conditions, rising healthcare costs and pressures on healthcare budgets. Drugs remain heavily scrutinized for cost containment. As a result, payers have been and continue to be more restrictive regarding the use of biopharmaceutical products and are scrutinizing the prices of these products while requiring a higher level of clinical evidence to support the benefits such products bring to patients and the broader healthcare system. For example, as discussed below, payers are increasingly using stricter utilization management criteria, such as prior authorization and step therapy, to contain or reduce costs. These pressures become intensified when our products become subject to competition, including from biosimilars.

In the United States, healthcare providers and other entities such as pharmacies and PBMs are reimbursed for covered services and products they deliver through both private-payer and government healthcare programs such as Medicare and Medicaid. We provide negotiated rebates or discounts to healthcare providers, private payers, government payers and PBMs. In addition, we are required to (i) provide rebates or discounts on our products that are reimbursed through certain government programs, including Medicare and Medicaid, and (ii) provide discounts to qualifying healthcare providers under the 340B Program. There has been an increase in the number of entities participating in the 340B Program and in the volume of our medicines dispensed through 340B channels. As a result, this expanded utilization of the 340B Program has had a negative impact on the Company's financial performance.

Both private and some government payers use formularies to manage access to and utilization of drugs. A drug's inclusion and favorable positioning on a formulary are essential to ensure patients have full access to a particular drug. Even when access is available, some patients abandon their prescriptions for economic reasons. Payers continue to institute cost reduction and containment measures that lower drug utilization and/or spending altogether and/or shift a greater portion of the costs to patients. Such measures include, but are not limited to, more-limited benefit plan designs, higher patient co-pays or coinsurance obligations, limitations on patients' use of commercial manufacturer co-pay payment assistance programs (including through co-pay accumulator adjustment or maximization programs), stricter utilization management criteria (such as prior authorization and step therapy) before a patient may get access to a drug, higher-tier formulary placement that increases the level of patient out-of-pocket costs and formulary exclusion, which effectively encourages patients and providers to seek alternative treatments or pay 100% of the cost of a drug. The use of such measures by PBMs and insurers has continued to intensify and has thereby limited Amgen product usage and sales. Furthermore, in the United States, the top six integrated health plans and PBMs controlled about 89% of all pharmacy prescriptions. As a result, PBMs and insurers have greater market power and negotiating leverage to mandate stricter utilization criteria and/or exclude drugs from their formularies in favor of competitor drugs or alternative treatments. In highly competitive treatment markets such as the markets for ENBREL, Otezla, Repatha and

Aimovig, PBMs are also able to exert negotiating leverage by requiring incremental rebates from manufacturers in order for them to gain and/or maintain their formulary position.

In addition to market actions taken by private and government payers in the United States, policy makers in both of the major U.S. political parties have supported policies to lower drug costs. See Item 1A. Risk Factors—*Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability.* For example, in 2022, the IRA was enacted and includes provisions requiring that beginning in 2026, mandatory price setting be introduced in Medicare for certain drugs paid for under Parts B and D, whereby manufacturers must accept a price established by the government or face penalties on all U.S. sales (starting with 10 drugs effective January 1, 2026, adding 15 in 2027 and 2028, and adding 20 in 2029 and subsequent years such that by 2031 approximately 100 drugs would be subject to such set prices). CMS has set Medicare Part D prices for ENBREL, effective January 1, 2026, and Otezla, effective January 1, 2027, in each case at significantly lower prices that are expected to negatively impact their profitability in Medicare. See Part IV—Note 13, Goodwill and other intangible assets, to the Consolidated Financial Statements for additional information related to the Otezla intangible asset impairment charges. Also under the IRA, starting on January 1, 2024, Medicare Part D was redesigned to cap beneficiary out-of-pocket costs and reduce Federal reinsurance in the catastrophic phase, increasing cost-sharing obligations for Part D plans and manufacturers, including by requiring manufacturer discounts. Further, the IRA created a mechanism for CMS to collect rebates from manufacturers if Medicare price increases outpace inflation. We began to accrue for rebate obligations on October 1, 2022 for Medicare Part D and on January 1, 2023 for Medicare Part B.

In addition to the IRA, other proposed U.S. policy actions focus on drug pricing, including the Most-Favored-Nations Prescription Drug Pricing Executive Order (MFN EO) that is aimed at using price benchmarks from other developed countries to set U.S. pricing targets, and the July MFN Letter that was delivered to a number of pharmaceutical companies, including Amgen. The July MFN Letter called for drug manufacturers to: 1) extend MFN pricing to Medicaid; 2) guarantee MFN pricing to Medicaid, Medicare and commercial payers on all newly launched drugs; 3) use future increased revenues from outside the U.S. to lower U.S. drug prices; and 4) participate in direct-to-patient models to provide MFN pricing for certain drugs. In December 2025, we announced that we are taking actions that satisfy the components outlined in the July MFN Letter, including the Administration's MFN pricing requests. We also announced the expansion of our direct-to-patient program, AmgenNow™, to include Aimovig and AMJEVITA, supplementing our existing offering of Repatha, all of which are available through AmgenNow™ and are expected to be available through TrumpRx following the platform's launch. See Item 1A. Risk Factors—*Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability.*

Other potential policies cover a wide range of areas, including allowing the importation of drugs from other countries; increasing transparency in drug pricing; using third-party value assessments to determine drug prices; referencing foreign prices; and changes to government rebate programs. For example, on January 5, 2024, the FDA authorized Florida to move forward with its importation program proposal, which excludes biologics. Further, at the state level, eight states (Colorado, Maine, Maryland, Minnesota, New Hampshire, New Jersey, Oregon and Washington) have enacted laws that establish PDABs to identify drugs that pose affordability challenges, and in four states (Colorado, Maryland, Minnesota and Washington) include authority for the state PDAB to set upper payment limits on certain drugs for in-state patients, payers and providers.

In many countries outside the United States, government-sponsored healthcare systems are the main payers for drugs and biologics. Governments and payers are using a range of tools to drive prices down, including mandatory price controls, international reference pricing, therapeutic-reference pricing, higher rebates, incentives for generics and biosimilars, and mandated price cuts. Pricing and reimbursement reforms under consideration in the United States—including proposals to expand government negotiation and increase payer leverage—may further amplify global pricing pressures, as many countries reference or react to U.S. pricing dynamics. Health technology assessment bodies increasingly evaluate the added benefit of new therapies, and these judgments shape reimbursement levels and coverage terms. Many countries also narrow coverage relative to product labels or impose volume caps to limit use. We expect governments to continue taking aggressive steps to reduce spending on drugs and biologics, and fiscal constraints may limit willingness to approve or provide access to innovative therapies and new technologies. Potential EU pharmaceutical legislation may also influence pricing and reimbursement frameworks, which could affect patient access.

The dynamics and developments discussed above create pressures on the pricing and potential usage of our products and on the industry. Given the diverse interests in play between payers, biopharmaceutical manufacturers, policy makers, healthcare providers and independent organizations, if and whether the parties involved can achieve alignment on the matters discussed above remain unclear, and the outcome of any such alignment is difficult to predict. We remain focused on pricing our products responsibly and delivering breakthrough treatments for unmet medical needs. Amgen is committed to working with the entire healthcare community to ensure continued innovation and to facilitate patient access to needed medicines. We do this by:

- investing billions of dollars annually in R&D;

- pricing our medicines to reflect the value they provide;
- developing more affordable therapeutic choices in the form of high-quality and reliably supplied biosimilars;
- partnering with payers to share risk and accountability for health outcomes;
- providing patient support and education programs;
- expanding patient access to our medicines, including through direct-to-patient channels;
- continuing to deliver a reliable supply of medicines through our additional investments in U.S. manufacturing;
- helping patients in financial need access our medicines; and
- working with policy makers, patients and other stakeholders to establish a sustainable healthcare system with access to affordable care and in which patients and their healthcare professionals are the primary decision makers.

See Item 1A. Risk Factors—*Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability* and Item 1A. Risk Factors—*Guidelines and recommendations published by various organizations can reduce the use of our products.*

Manufacturing, Distribution and Raw Materials

Manufacturing

We believe we are a leader in the manufacture of biologics and that our manufacturing capabilities represent a competitive advantage. The products we manufacture consist of both biologics and small molecule drugs. The majority of our products are biologics that are produced in living cells and that are inherently complex due to naturally occurring molecular variations. Highly specialized knowledge and extensive process and product characterization are required to transform laboratory-scale processes into reproducible commercial manufacturing processes. Further, our expertise in the manufacture of biologics has positioned us well for leadership in the global biosimilars market. For additional information regarding manufacturing facilities, see Item 2. Properties.

We are expanding our manufacturing capacity and incorporating state-of-the-art technologies to allow us to optimize our manufacturing network and mitigate risks while working to continue to ensure adequate supply of our products to patients worldwide. See *Manufacturing Initiatives* below for additional discussion of our manufacturing expansion efforts. Our new state-of-the-art biomanufacturing facilities, including our facility in North Carolina and FDA-approved facility in Ohio, have been constructed at a lower cost and with greater speed as compared to traditional facilities. For example, our facility in North Carolina has equipment that is portable and smaller, which provides greater flexibility and speed in the manufacture of different medicines simultaneously, allowing us to respond to changing demands for our medicines with increased scale and agility. Furthermore, such state-of-the-art facilities incorporate multiple innovative technologies, automation solutions and environmental sustainability, thus requiring smaller manufacturing footprints and offering greater environmental benefits, including reduced consumption of water and energy and lower levels of carbon emissions. For example, our facility in North Carolina and our FDA-approved facility in Ohio contain many examples of environmental commitments, including on-site photovoltaic renewable energy generation at both sites. We expect our North Carolina facility's carbon footprint, water usage and waste disposed to be substantially lower than that of a traditional drug substance manufacturing facility. Similarly, we expect lower carbon footprint, water usage and waste disposed per unit at our Ohio facility as compared to traditional packaging and assembly facilities.

Our internal manufacturing network has commercial production capabilities for bulk manufacturing, formulation, fill, finish, tableting and final device assembly. These activities are performed within the United States, including in the U.S. territory of Puerto Rico, Rhode Island, Ohio and California facilities, as well as internationally in our Ireland, Netherlands and Singapore facilities. In addition, we use third-party contract manufacturers to supplement the capacity or capability of our commercial manufacturing network.

To support our clinical trials, we manufacture product candidates primarily at our California facilities. We also use third-party contract manufacturers, including contract manufacturers that were added to our clinical manufacturing network from the Horizon acquisition, to supplement the capacity or capability of our overall clinical manufacturing network.

See Item 1A. Risk Factors for a discussion of the factors that could adversely impact our manufacturing expansion operations and the global supply of our products.

Distribution

We operate distribution centers in the U.S. territory of Puerto Rico, Kentucky, California and the Netherlands for worldwide distribution of the majority of our commercial and clinical products. We also use third-party distributors to supplement distribution of our products worldwide.

Other

In addition to the manufacturing and distribution activities noted above, each of our manufacturing locations includes key manufacturing support functions such as quality control, process development, engineering, procurement, production scheduling and warehousing. Certain of those manufacturing and distribution activities are highly regulated by the FDA as well as international regulatory agencies. See Government Regulation—*Regulation in the United States—Regulation of Manufacturing Standards*.

Manufacturing Initiatives

As discussed above, we are expanding our capacity and advancing new innovations to support anticipated patient demand for our current and future products.

In January 2025, we opened our Holly Springs, North Carolina site, a cutting-edge drug substance facility. Upon FDA approval, this facility will increase our biologics manufacturing capacity. Also in January 2025, we broke ground on a second drug substance manufacturing facility at this location. This second facility will incorporate state-of-the-art technologies and sustainable practices. The result is a more flexible and efficient facility, with a smaller physical footprint, reduced consumption of water and energy, and lower levels of carbon emissions than a traditional facility.

Additionally, in 2025 we announced expansions of our manufacturing network in New Albany, Ohio and the U.S. territory of Puerto Rico, further strengthening our U.S.-based manufacturing capabilities. Our Ohio expansion builds on our existing biomanufacturing facility, which received FDA licensure for commercial production in January 2024 and supports final product assembly and packaging using state-of-the-art technologies and automation.

These expansion initiatives further enhance our manufacturing network to support reliable and efficient supply of our medicines to patients worldwide. We continue evaluating our manufacturing network to ensure we maintain appropriate levels of operational flexibility and risk management needed to support our global operations.

Numerous tariffs and trade protection measures have been proposed and, in some cases, implemented by the United States and other countries. We continue to evaluate our global supply chains, sourcing strategies and manufacturing network and to implement measures to minimize the impact of existing and potential future tariffs. See Item 1A. Risk factors—*Our sales and operations are subject to the risks of doing business internationally, including in new or emerging markets*, and Part II—Item 7. Management’s Discussion and Analysis of Financial Condition and Results of Operations—Overview.

See Item 1A. Risk Factors—*Manufacturing difficulties, disruptions or delays could limit supply of our products and limit our product sales*.

Raw Materials and Medical Devices

Certain raw materials, medical devices (including companion diagnostics) and components necessary for the commercial and/or clinical manufacturing of our products are provided by and are the proprietary products of unaffiliated third-party suppliers, certain of which may be our only source for such materials. We work to manage the risk associated with such sole suppliers by means of inventory management, relationship management and evaluation of alternative sources when feasible. We also monitor the financial condition and manufacturing quality and compliance of key suppliers and their ability to supply our needs. See Item 1A. Risk Factors—*We rely on third-party suppliers for certain of our raw materials, medical devices and components*.

We perform various procedures to help authenticate the sources of raw materials, including intermediary materials used in the manufacture of our products; the procedures are incorporated into the manufacturing processes we and our third-party contract manufacturers perform and include verification of country of origin.

To better ensure supply, Amgen has a risk mitigation strategy that uses a combination of methods, including multiple sources or backup inventory of critical raw materials. As part of our ongoing business continuity efforts, we continue to closely monitor our inventory levels and have taken additional measures to mitigate against raw material supply interruption. We also continue to assess the potential impacts of existing and proposed tariffs on our raw material and medical device supply chains and implement sourcing and inventory strategies intended to minimize the impacts from existing and potential future tariffs. See

Item 1A. Risk Factors for a discussion of the factors that could adversely impact our manufacturing operations and the global supply of our products.

Government Regulation

Regulation by government authorities in the United States and other countries is a significant factor in the production and marketing of our products and our ongoing R&D activities. To clinically test, manufacture and market products for therapeutic use, we must satisfy mandatory procedures and safety and effectiveness standards established by various regulatory bodies. Compliance with these standards is complex, and failure to comply with any of these standards can result in significant implications. See Item 1A. Risk Factors for a discussion of factors, including global regulatory implications, that can adversely impact our development and marketing of commercial products.

Regulation in the United States

In the United States, the Public Health Service Act; the FDCA; and the regulations promulgated thereunder as well as other federal and state statutes and regulations govern, among other things, the production, research, development, testing, manufacture, quality control, labeling, storage, record keeping, approval, advertising, promotion and distribution of our products in addition to the reporting of certain payments and other transfers of value to healthcare professionals and teaching hospitals.

Clinical Development and Product Approval. Drug development in our industry is complex, challenging and risky, and failure rates are high. Product development cycles are typically very long—approximately 10 to 15 years from discovery to market. A potential new medicine must undergo many years of preclinical and clinical testing to establish its safety and efficacy for use in humans at appropriate dosing levels and with an acceptable risk–benefit profile. We continue to work toward reducing cycle times by applying our expertise in human genetics and innovation in technology, clinical trials and real-world evidence. See Research and Development and Selected Product Candidates section below.

After laboratory analysis and preclinical testing in animals, we file an IND with the FDA to begin human testing. Typically, we undertake an FDA-designated three-phase human clinical testing program.

- In Phase 1, we conduct small clinical trials to investigate the safety and proper dose ranges of our product candidates in a small number of human subjects.
- In Phase 2, we conduct clinical trials to investigate side-effect profiles and the efficacy of our product candidates in a patient population larger than Phase 1 but still relatively small, who have the disease or condition under study.
- In Phase 3, we conduct clinical trials to investigate the short- and long-term safety and efficacy of our product candidates, compared to commonly used treatments, in a large number of patients who have the disease or condition under study.

The FDA monitors the progress of each trial conducted under an IND and may, at its discretion, reevaluate, alter, suspend or terminate the testing based on data accumulated to that point and the FDA's risk–benefit assessment with regard to the patients enrolled in the trial. The results of preclinical and clinical trials are submitted to the FDA in the form of either a BLA for biologic products or a New Drug Application for small molecule products. We are not permitted to market or promote a new product until the FDA has approved our marketing application.

Approval of Biosimilars. The Affordable Care Act authorized the FDA to approve biosimilars via a separate, abbreviated pathway. The pathway allows sponsors of a biosimilar to seek and obtain regulatory approval based in part on the nonclinical-trial and clinical-trial data of an originator product to which the biosimilar has been demonstrated to be “highly similar” and to have no clinically meaningful differences with regard to safety, purity and potency. The relevance of demonstrating “similarity” is that in many cases, biosimilars can be brought to market without conducting the full suite of clinical trials typically required of originators, because risk–benefit has previously been established. To preserve incentives for future innovation, the law establishes a period of exclusivity for originators' products, which in general prohibits biosimilars from gaining FDA approval based in part on reliance on or reference to the originator's data in their application to the FDA for 12 years after initial FDA approval of the originator product. The law does not change the duration of patents granted on biologic products. As part of the implementation of the abbreviated approval pathway for biosimilars, the FDA released a number of guidance documents, some of which remain in draft form. See Item 1A. Risk Factors—*We currently face competition from biosimilars and generics and expect to face increasing competition from biosimilars and generics in the future.*

Regulation of Product Marketing and Promotion. The FDA regulates the marketing and promotion of drug products. Our product promotions for approved product indications must comply with the statutory standards of the FDCA and the FDA's implemented regulations and guidance. The FDA's review of marketing and promotional activities encompasses but is not limited to direct-to-patient advertising, healthcare-provider-directed advertising and promotion, sales representative communications to healthcare professionals, promotional programming and promotional activities involving electronic media. The FDA may also review industry-sponsored scientific and educational activities that make representations regarding product safety or efficacy in a promotional context. The FDA may take enforcement action against a company for violations of the FDA's advertising and labeling laws and regulations. Enforcement action may include product seizures, injunctions, civil or criminal penalties or regulatory letters, which may require corrective advertising or other corrective communications to healthcare professionals. Failure to comply with the FDA's regulations also can result in adverse publicity or increased scrutiny of company activities by the U.S. Congress or other legislators. Additionally, as described below, such failure may lead to additional liability under U.S. healthcare fraud and abuse laws.

Regulation of Manufacturing Standards. The FDA regulates and inspects the equipment, facilities, laboratories and processes used in the manufacturing and testing of products prior to granting approval to market products. If, after receiving approval from the FDA, we make a material change in manufacturing equipment, location or process, additional regulatory review may be required. We also must adhere to current Good Manufacturing Practice regulations and product-specific regulations enforced by the FDA through its facilities inspection program. The FDA conducts regular, periodic visits to reinspect our equipment, facilities, laboratories and processes following an initial approval.

Regulation of Combination Products. Combination products are defined by the FDA as products composed of two or more regulated components (e.g., a biologic and/or drug and a device). Biologics/drugs and devices each have their own regulatory requirements, and combination products may have additional requirements. A number of our marketed products meet this definition and are regulated under this framework, and we expect that a number of our pipeline product candidates will be evaluated for regulatory approval under this framework as well.

Regulation of Orphan Medicines. Orphan drugs are defined by the FDA as products intended to treat a rare disease or condition that affects less than 200,000 persons in the United States. A company must request orphan drug designation prior to filing, and if the product is ultimately approved for the designated rare disease or condition, it may qualify for seven years of orphan drug exclusivity during which the FDA will not approve another sponsor's marketing application for the same drug for the same indication. Orphan drug exclusivity will not bar approval of another medicine for the same indication if it is shown to be clinically superior. In the United States, a number of our products, including products such as BLINCYTO and UPLIZNA, have orphan drug exclusivity under this framework for specific indications.

Regulation Outside the United States

In EU countries as well as in the United Kingdom, Switzerland, Canada, Australia and Japan, regulatory requirements and approval processes are similar in principle to those in the United States.

In the EU, there are currently two potential tracks for seeking marketing approval for a product not authorized in any EU member state: a decentralized procedure and a centralized procedure. In the *decentralized procedure*, identical applications for marketing authorization are submitted simultaneously to the national regulatory agencies. Regulatory review is led by one member state (the reference-member state), and its assessment—based on safety, quality and efficacy—is reviewed and approved (assuming there are no concerns that the product poses a serious risk to public health) by the other member states from which the applicant is seeking approval (the concerned-member states). The decentralized procedure leads to a series of single national approvals in all relevant countries. In the *centralized procedure*, which is required of all products derived from biotechnology, a company submits a single MAA to the EMA, which conducts an evaluation of the dossier, drawing upon its scientific resources across Europe. If the drug product is proven to fulfill requirements for quality, safety and efficacy, the EMA's CHMP adopts a positive opinion, which is transmitted to the EC for final decision on granting of the marketing authorization. Even though the EC generally follows the CHMP's opinion, it is not bound to do so. Subsequent commercialization is enabled by country-by-country reimbursement approval.

In the EU, biosimilars are approved under a specialized pathway of the centralized procedure. As with the U.S. pathway, an applicant seeks and obtains regulatory approval for a biosimilar once the data exclusivity period for the original reference product has expired, relying, in part, on the data submitted for the originator product together with data evidencing that the biosimilar is "highly similar" with regard to quality, safety and efficacy to the original reference product authorized in the European Economic Area. See Item 1A. Risk Factors—*We currently face competition from biosimilars and generics and expect to face increasing competition from biosimilars and generics in the future.*

In the EU, Regulation (EC) No 141/2000, as implemented by Regulation (EC) No. 847/2000, provides that a medicine can be designated as an orphan medicinal product by the EC if its sponsor can establish that: (i) the product is intended for the diagnosis, prevention or treatment of life-threatening or chronically debilitating conditions; (ii) either (a) such conditions affect not more than 5 in 10,000 persons in the EU when the application is made, or (b) the product without the benefits derived from orphan status, would not generate sufficient return in the EU to justify the necessary investment in developing the medicinal product; and (iii) there exists no satisfactory authorized method of diagnosis, prevention, or treatment of the condition that has been authorized in the EU, or even if such method exists, the product will be of significant benefit to those affected by that condition. An application for the designation of a medicinal product as an orphan medicinal product may be submitted at any stage of development of the medicinal product but before the filing of an MAA. A marketing authorization for an orphan medicinal product may only include indications designated as orphan. For non-orphan indications treated with the same active pharmaceutical ingredient, a separate marketing authorization has to be sought. Approved orphan drugs in the EU receive 10 years of market exclusivity for the approved indication in all EU member states. We currently have orphan medicinal product designation for BLINCYTO in the EU and may seek medicinal product designation for a number of our products in the future.

Other countries such as those in Latin America and the Middle East have review processes and data requirements similar to those of the EU and in some cases can rely on prior marketing approval from U.S. or EU regulatory authorities. The regulatory process in these countries may include manufacturing/testing facility inspections, testing of drug product upon importation and other domestic requirements.

In Asia Pacific, a number of countries such as China, Japan, South Korea and Taiwan may require local clinical-trial data for bridging purposes as part of the drug registration process in addition to global clinical trials, which can add to overall drug development and registration timelines. In many Asian markets, registration timelines may be influenced by prior marketing approvals in reference jurisdictions (e.g. the United States or the EU). In some markets in Asia, such as China, Indonesia and Thailand, regulatory timelines can be more variable. The regulatory process may also include manufacturing/testing facility inspections, testing of drug product upon importation and other domestic requirements. Countries such as Australia and Japan have more mature regulatory systems which may allow for submissions under comparatively more predictable time frames. With regard to biosimilars, several of these countries have pathways to register biosimilars (e.g., Australia, India, Singapore, South Korea and Taiwan), and biosimilar products in market (e.g., Australia, Japan, South Korea and Taiwan).

In some countries, such as Japan and those in the EU, medical devices may be subject to regulatory regimes whereby manufacturers must establish that their medical devices conform to essential requirements set out in the law for the particular device category. For example, in the EU, with limited exceptions, medical devices placed on the market must bear the Conformité Européenne marking to indicate their conformity with legal requirements.

Post-approval Phase

After approval, we continue to monitor adverse events and product complaints reported following the use of our products through routine postmarketing surveillance and studies when applicable. We report such events to the appropriate regulatory agencies as required by local regulations for individual cases and aggregate reports. We proactively monitor (according to good pharmacovigilance practices) and ensure the implementation of signal detection, assessment and the communication of adverse events that may be associated with the use of our products. We also proactively monitor product complaints through our quality systems, which includes assessing our drug delivery devices for device complaints, adverse events and malfunctions. We may also be required by regulatory agencies to conduct further clinical trials on our marketed products as a condition of their approval or to provide additional information on safety and efficacy. Health regulators, including the FDA, have authority to mandate labeling changes to products at any point based on new safety information or as part of an evolving label change to a particular class of products.

Health regulators, including the FDA, also have authority both before and after approval to require that a company implement a risk management program for a product to ensure that the benefits of the drug outweigh the risks. Each risk management program is unique and varies depending on the specific factors required. In the United States, such a risk management program is known as a REMS, and we currently have REMSs for Prolia, Nplate and BLINCYTO.

Other Regulation

We are also subject to various laws pertaining to healthcare fraud and abuse, including antikickback laws and false-claims laws. Antikickback laws make it illegal to solicit, offer, receive or pay any remuneration in exchange for or to induce the referral of business, including the purchase or prescribing of a particular drug that is reimbursed by a state or federal program. False-claims laws prohibit knowingly and willingly presenting, or causing to be presented, for payment to third-party payers (including Medicare and Medicaid) any claims for reimbursed drugs or services that are false or fraudulent, claims for items or services not provided as claimed, or claims for medically unnecessary items or services. Violations of fraud and abuse laws may be punishable by criminal and/or civil sanctions, including fines and civil monetary penalties, as well as by the possibility

of exclusion from federal healthcare programs (including Medicare and Medicaid). Liability under false-claims laws may also arise when violation of certain laws or regulations related to the underlying product (e.g., a violation regarding improper promotional activity or unlawful payments) contributes to the submission of a false claim. See Item 1A. Risk Factors—*Our business may be affected by litigation and government investigations.*

The FCPA prohibits U.S. corporations and their representatives from offering, promising, authorizing or making payments to any foreign government official, government staff member, political party or political candidate in an attempt to obtain or retain business abroad. The scope of the FCPA arguably includes interactions with certain healthcare professionals in many countries. Other countries have enacted similar anticorruption laws and/or regulations. Failure by our employees, agents, contractors, vendors, licensees, partners or collaborators to comply with the FCPA and other anticorruption laws and/or regulations could result in significant civil or criminal penalties.

We are subject to various laws and regulations globally with regard to privacy and data protection. These laws and regulations involve the collection, storage, handling, use, disclosure, transfer and security of personal data. The legislative and regulatory environments regarding privacy and data protection are continually evolving and developing because these issues are subjects of increasing amounts of attention in countries globally. For example, we are subject to the EU's GDPR, which became effective on May 25, 2018; the CCPA, which became effective on January 1, 2020; the California Privacy Rights Act of 2020, which amended the CCPA and became effective on January 1, 2023; and China's Personal Information Protection Law, which became effective on November 1, 2021. Other jurisdictions where we operate have enacted or proposed similar legislation and/or regulations, such as consumer privacy laws that went into effect in Virginia, Colorado, Utah, Connecticut and Florida in 2023 and Oregon, Texas and Montana in 2024. Consumer privacy laws were also passed in other states, including Iowa, Delaware, New Hampshire, Nebraska, New Jersey, Tennessee, Minnesota, Maryland, Indiana, Kentucky and Rhode Island, and became effective or are scheduled to become effective in 2025 through 2026. In April 2023, a new type of state privacy law focused on protection of consumer health data emerged in Washington with the enactment of the My Health My Data Act, with similar legislation passed subsequently in Nevada. Both these new consumer health privacy laws became effective on March 31, 2024. Further, in 2024, the EU Artificial Intelligence (AI) Act, formally known as Regulation (EU) 2024/1689, was passed into law. Certain provisions of this regulation, such as transparency obligations and governance structures, became effective in February and August 2025, and the regulation will become fully effective on August 2, 2026. The regulation establishes a risk-based framework governing the development, deployment and use of AI systems across the EU. High-risk AI systems are subject to stringent requirements such as mandatory risk assessments, technical documentation, bias mitigation and explainability standards. Non-compliance with these current and future laws could result in significant penalties.

Our business has been and will continue to be subject to various other U.S. and foreign laws, rules and regulations, including provisions of the IRA and OB3. See Reimbursement section above.

Research and Development and Selected Product Candidates

We focus our R&D on novel human therapeutics for the treatment of serious illness. We capitalize on our strengths in human genetics, novel biology and protein engineering. We leverage our biologic expertise and seek to choose the optimal modality for a drug target and disease, and we use cutting-edge science and technology to study subtle biological mechanisms in search of therapies that will improve the lives of those who suffer from diseases. See Government Regulation—*Clinical Development and Product Approval* section above.

Our discovery research programs may therefore yield targets that lead to the development of human therapeutics delivered as large molecules, small molecules, other combination modalities or new modalities. We have increasingly focused our efforts on human genetics when possible to enhance the likelihood of success. We have major R&D centers in the United States in Thousand Oaks and San Francisco, California; Iceland; and the United Kingdom, as well as smaller research centers and development facilities globally. Additionally, in 2025, we broke ground on a new state-of-the-art R&D facility in Thousand Oaks, California, to further enhance collaboration and innovation across R&D and process development activities. See Item 2. Properties.

Our clinical trial activities are conducted by both our internal staff and third-party contract clinical trial service providers. To increase the number and diversity of patients available for enrollment in our clinical trials and to meet evolving regulatory expectations, we have opened clinical sites and will continue opening clinical sites and enrolling patients in a number of geographic locations. With regard to our clinical trial activities, we are continuously monitoring the possible impacts from health-related events, geopolitical conflicts and natural disasters. We are working to mitigate effects on future study enrollment in our clinical trials; and we are evaluating the impact in all relevant countries. We remain focused on supporting our active clinical sites in their providing care for patients and in our providing investigational drug supply. See Government Regulation—*Regulation in the United States—Clinical Development and Product Approval* for a discussion of government regulation over clinical development. Also see Item 1A. Risk Factors—*We must conduct clinical trials in humans before we commercialize and sell any of our product candidates or existing products for new indications.*

For the years ended December 31, 2025, 2024 and 2023, our R&D expenses were \$7.3 billion, \$6.0 billion and \$4.8 billion, respectively. We expect to continue allocating significant resources to our R&D activities.

Some of our competitors are actively engaged in R&D in areas in which we have products or in which we are developing product candidates or new indications for existing products. For example, we compete with other clinical trials for eligible patients, which may limit the number of available patients who meet the criteria for certain clinical trials. The competitive marketplace for our product candidates is greatly dependent on the timing of entry into the market. Early entry may have important advantages in gaining product acceptance, thereby contributing to a product's eventual success and profitability. Accordingly, we expect that in some cases, the relative speed with which we can develop products, complete clinical testing, receive regulatory approval and supply commercial quantities of a product to the market will be important to our competitive position.

In addition to product candidates and marketed products generated from our internal R&D efforts, we acquire companies, acquire and license certain product and R&D technology rights and establish R&D arrangements with third parties to enhance our strategic position within our industry by strengthening and diversifying our R&D capabilities, product pipeline and marketed product base. In pursuing these R&D arrangements and licensing or acquisition activities, we face competition from other pharmaceutical and biotechnology companies that also seek to license or acquire technologies, product candidates or marketed products from those entities performing the R&D.

The following table shows a selection of certain of our product candidates by phase of development in our therapeutic areas of focus as of February 3, 2026, unless otherwise indicated. The information in this section does not include, among other things, other, nonregistrational clinical trials that we may conduct for purposes other than for submission to regulatory agencies for their approval of a new product indication.

We may conduct nonregistrational clinical trials for various reasons, including to evaluate real-world outcomes or to collect additional safety information with regard to the use of products.

	Investigational indications (programs)
Phase 3	
BLINCYTO	Ph-negative B-ALL
Dazodalibep	Sjögren's disease
IMDELLTRA/IMDYLLTRA	Small cell lung cancer
LUMAKRAS/LUMYKRAS	Metastatic colorectal cancer
	Non-small cell lung cancer
MariTide	Chronic weight management
	Cardiovascular disease
	Heart failure
	Obstructive sleep apnea
Nplate	Chemotherapy-induced thrombocytopenia
Olpasiran	Cardiovascular disease
Pegloticase	Subcutaneous administration for uncontrolled gout
Repatha	Cardiovascular disease
Rocatinlimab ⁽¹⁾	Moderate-to-severe atopic dermatitis
	Prurigo nodularis
TEPEZZA	Subcutaneous administration for TED
	Chronic/low clinical activity score TED in Japan
TEZSPIRE	Chronic obstructive pulmonary disease
	Eosinophilic esophagitis
Xaluritamig	Metastatic castrate resistant prostate cancer
ABP 206	Investigational biosimilar to OPDIVO® (nivolumab)
ABP 234	Investigational biosimilar to KEYTRUDA® (pembrolizumab)
ABP 692	Investigational biosimilar to OCREVUS® (ocrelizumab)

	Investigational indications (programs)
Phase 2	
Blinatumomab	Refractory rheumatoid arthritis Systemic lupus erythematosus with and without nephritis
Daxdilimab	Dermatomyositis and anti-synthetase inflammatory myositis Discoid lupus erythematosus
Inebilizumab	Systemic lupus erythematosus with nephritis
MariTide	Type 2 diabetes
Rocatinlimab ⁽¹⁾	Moderate-to-severe asthma
AMG 104	Asthma
AMG 193	Non-small cell lung cancer
AMG 329	Sjögren's disease
AMG 732	TED
Phase 1	
Xaluritamig	Ewing Sarcoma
AMG 193	Other tumors
AMG 305	Solid tumors
AMG 355	Solid tumors
AMG 410	Solid tumors
AMG 513	Obesity
AMG 691	Asthma

⁽¹⁾ See Significant Developments for additional information regarding the termination of our collaboration agreement with Kyowa Kirin.

Phase 3 Clinical trials investigate the short- and long-term safety and efficacy of our product candidates, compared to commonly used treatments, in a large number of patients who have the disease or condition under study.

Phase 2 Clinical trials investigate side-effect profiles and efficacy of product candidates in a larger patient population than Phase 1, but still relatively small, who have the disease or condition under study.

Phase 1 Clinical trials investigate the safety and proper dose ranges of product candidates usually in a small number of human subjects.

Phase 3 Product Candidate Program Changes

As of February 4, 2025, we had 25 Phase 3 programs studied in investigational indications. As of February 3, 2026, we had 23 Phase 3 programs being studied in investigational indications, as six programs initiated Phase 3 studies, five programs received regulatory approval, one program was stopped, one program was no longer being pursued and one program concluded. These changes are set forth in the following table.

Molecule	Investigational indications (programs)	Program changes
AMJEVITA	Interchangeability study	Approved by the FDA
Bemarituzumab	Gastric and gastroesophageal junction cancer	Stopped
EVENTITY	Male osteoporosis	No longer pursued
MariTide	Chronic weight management	Initiated two global Phase 3 studies
	Cardiovascular disease	Initiated global Phase 3 study
	Heart failure	Initiated global Phase 3 study
	Obstructive sleep apnea	Initiated two global Phase 3 studies
Otezla	Palmoplantar pustulosis	Approved in Japan
Pegloticase	Subcutaneous administration for uncontrolled gout	Initiated Phase 3 study
TEZSPIRE	Chronic obstructive pulmonary disease	Initiated Phase 3 study
	Chronic rhinosinusitis with nasal polyps	Approved by the FDA and the EC
	Severe asthma	Concluded
UPLIZNA	IgG4-RD	Approved by the FDA and the EC
	Generalized myasthenia gravis	Approved by the FDA

Phase 3 Product Candidate Patent Information

The following table describes our composition-of-matter patents that have been issued thus far for our product candidates in Phase 3 development that have yet to be approved for any indication in the United States or the EU. Patents for products already approved for one or more indications in the United States or the EU but that are currently undergoing Phase 3 clinical trials for additional indications have been previously described. See Marketing, Distribution and Selected Marketed Products—*Patents*.

Molecule	Territory	General subject matter	Estimated expiration ⁽¹⁾
Dazodalibep	U.S.	Polypeptides	2034
	Europe	Polypeptides	2032
MariTide	U.S.	Antibody-peptide conjugates	2038
	Europe	Antibody-peptide conjugates	2038
Olpasiran	U.S.	Compounds	2036
	Europe	Compounds	2036
Rocatinlimab ⁽²⁾	U.S.	Polypeptides	2028
	Europe	Polypeptides	2026
Xaluritamig	U.S.	Polypeptides	2039

⁽¹⁾ Patent expiration estimates are based on issued patents, which may be challenged, invalidated or circumvented by competitors. The estimates do not include any term adjustments, extensions or supplemental protection certificates that may be obtained in the future and thereby extend these dates. Corresponding patent applications are pending in other jurisdictions. Additional patents may be filed or issued and may provide additional exclusivity for the product candidate or its use. In addition to patent exclusivity, the product candidates may be protected by regulatory exclusivities upon approval in some countries. For example, new chemical entities would receive a five year exclusivity period and new molecular entities would receive a 12 year exclusivity period in the United States, whereas new chemical and molecular entities would receive a 10 year exclusivity period in Europe.

⁽²⁾ See Significant Developments for additional information regarding the termination of our collaboration agreement with Kyowa Kirin.

Phases 2 and 3 Program Descriptions

The following provides additional information about selected products and product candidates that have advanced into human clinical trials.

Blinatumomab

Blinatumomab is an anti-CD19 x anti-CD3 BiTE[®] molecule. It is being investigated for the treatment of both refractory rheumatoid arthritis and systemic lupus erythematosus with and without nephritis.

BLINCYTO

BLINCYTO is an anti-CD19 x anti-CD3 BiTE[®] molecule. It is being investigated for the treatment of newly diagnosed adults with B-ALL.

Daxilimab

Daxilimab is a fully human monoclonal antibody against ILT7 that depletes certain dendritic cells. It is being investigated for the treatment of both dermatomyositis and anti-synthetase inflammatory myositis and discoid lupus erythematosus.

Dazodalibep

Dazodalibep is a fusion protein binding CD40L on T cells, blocking their interaction with CD40-expressing B cells. It is being investigated for the treatment of Sjögren's disease.

IMDELLTRA/IMDYLLTRA

IMDELLTRA/IMDYLLTRA is an anti-DLL3 x anti-CD3 BiTE[®] molecule. It is being investigated for the treatment of small cell lung cancer. See Significant Developments for additional information regarding clinical trial updates.

Inebilizumab

Inebilizumab is a humanized, affinity-optimized, afucosylated IgG1 kappa (IgG1κ) monoclonal antibody that binds to the B cell-specific surface antigen CD19. It is being investigated as a treatment for patients with systemic lupus erythematosus with nephritis.

LUMAKRAS/LUMYKRAS

LUMAKRAS/LUMYKRAS is a KRAS^{G12C} small molecule inhibitor. It is being investigated in colorectal cancer, NSCLC and other solid tumor cancers.

Maridebart cafraglutide

MariTide is a differentiated antibody-peptide conjugate that activates the GLP-1 receptor and antagonizes the GIPR. It is being investigated for the treatment of chronic weight management, cardiovascular disease, heart failure, obstructive sleep apnea and Type 2 diabetes. See Significant Developments for additional information regarding clinical trial updates.

Nplate

Nplate is a thrombopoietin receptor agonist (TPO-RA). It is being investigated for the treatment of chemotherapy-induced thrombocytopenia (CIT).

Olpasiran

Olpasiran is a small interfering RNA (siRNA) that lowers lipoprotein(a) (Lp(a)). It is being investigated for the treatment of atherosclerotic cardiovascular disease (ASCVD).

Pegloticase

Pegloticase is a PEGylated uricase approved for the treatment of uncontrolled gout by intravenous administration. It is being investigated for subcutaneous administration when co-administered with methotrexate.

Repatha

Repatha is a human monoclonal antibody that inhibits PCSK9. See Significant Developments for additional information regarding clinical trial updates.

Rocatinlimab

Rocatinlimab is a monoclonal antibody that inhibits OX-40. It is being investigated in Phase 3 studies for the treatment of moderate-to-severe atopic dermatitis and prurigo nodularis. It is also being investigated in a Phase 2 study for the treatment of moderate-to-severe asthma. See Significant Developments for additional information regarding the termination of our collaboration agreement with Kyowa Kirin.

TEPEZZA

TEPEZZA is a monoclonal antibody against IGF-1R. It is being investigated for subcutaneous administration for the treatment of TED and chronic/low clinical activity score TED in Japan.

TEZSPIRE

TEZSPIRE is a human monoclonal antibody that inhibits the action of thymic stromal lymphopoietin. It is being investigated for the treatment of chronic obstructive pulmonary disease and eosinophilic esophagitis. TEZSPIRE is being developed in collaboration with AstraZeneca.

Xaluritamig

Xaluritamig is an anti-six transmembrane epithelial antigen of the prostate 1 (STEAP1) x anti-CD3 XmAb[®] BiTE[®] molecule. It is being investigated for the treatment of prostate cancer.

ABP 206

ABP 206, a biosimilar candidate to OPDIVO[®], is a monoclonal antibody that binds to the receptor protein called programmed death protein 1 (PD-1).

ABP 234

ABP 234, a biosimilar candidate to KEYTRUDA[®], is a monoclonal antibody that binds to the receptor protein PD-1. It is being investigated in a Phase 3 study for biosimilarity to KEYTRUDA[®]. The reference-product primary condition is NSCLC.

ABP 692

ABP 692, a biosimilar candidate to OCREVUS[®], is a monoclonal antibody that binds to CD20, which is a protein found on the surface of B-cells.

AMG 104

AMG 104 is a human anti-TSLP Fab. It is being investigated for the treatment of asthma and is being developed in collaboration with AstraZeneca.

AMG 193

AMG 193 is a small molecule methylthioadenosine (MTA) cooperative protein arginine methyltransferase 5 (PRMT5) inhibitor. It is being investigated for the treatment of NSCLC.

AMG 329

AMG 329 is a fully human monoclonal antibody that binds and neutralizes the function of the FLT3-ligand, thereby reducing both conventional and plasmacytoid dendritic cells. It is being investigated for the treatment of Sjögren's disease.

AMG 732

AMG 732 is a monoclonal antibody against IGF-1R. It is being investigated for the treatment of TED.

Business Relationships

From time to time, we enter into business relationships, including joint ventures and collaborative arrangements, for the R&D, manufacture and/or commercialization of products and/or product candidates. In addition, we acquire product and R&D technology rights and establish R&D collaborations with third parties to enhance our strategic position within our industry by strengthening and diversifying our R&D capabilities, product pipeline and marketed-product base. These arrangements generally provide for nonrefundable upfront license fees, development and commercial-performance milestone payments, cost sharing, royalties and/or profit sharing. The activities under these collaboration agreements are performed with no guarantee of either technological or commercial success, and each is unique in nature.

Trade secret protection for our unpatented confidential and proprietary information is important to us. To protect our trade secrets, we generally require counterparties to execute confidentiality agreements upon commencement of a business relationship with us. However, others could either develop independently the same or similar information or unlawfully obtain access to our information.

AstraZeneca plc

We are in a collaboration with AstraZeneca for the development and commercialization of TEZSPIRE.¹ Under our collaboration, both companies share global costs, profits and losses equally after payment by AstraZeneca of a mid-single-digit royalty to Amgen. AstraZeneca leads global development. In North America, Amgen, as the principal, recognizes product sales of TEZSPIRE in the United States, and AstraZeneca, as the principal, recognizes product sales of TEZSPIRE in Canada. AstraZeneca leads commercialization for TEZSPIRE outside North America. Amgen manufactures and supplies TEZSPIRE worldwide.

UCB

We are in a collaboration with UCB for the development and commercialization of EVENITY. Under our collaboration, UCB has rights to lead commercialization for EVENITY in most countries in Europe. Amgen, as the principal, leads commercialization for EVENITY and recognizes product sales in all other territories, including the United States. Global development costs and commercialization profits and losses related to the collaboration are shared equally. Amgen manufactures and supplies EVENITY worldwide.

BeOne Medicines Ltd.

In January 2020, we acquired an equity stake in BeOne for approximately \$2.8 billion in cash as part of a collaboration agreement to expand our oncology presence in China. For additional information regarding our equity investment in BeOne, see Part IV—Note 10, Investments, to the Consolidated Financial Statements. Under the collaboration, BeOne began selling XGEVA in 2020, BLINCYTO in 2021 and KYPROLIS in 2022 in China, and Amgen shares profits and losses equally during the product-specific commercialization periods. Following entry into an amendment to this collaboration by Amgen and BeOne on November 11, 2025, BeOne will retain the right to commercialize XGEVA, BLINCYTO and KYPROLIS in China for so long as such products are sold in China. Amgen manufactures and supplies the collaboration products to BeOne.

In addition, we jointly develop a portion of our oncology portfolio with BeOne, which shares in global R&D costs by providing cash and development services of up to \$1.25 billion. Upon regulatory approval, BeOne will assume commercialization rights in China for a specified period, and Amgen and BeOne will share profits and losses equally until certain of these product rights revert to Amgen. Upon return of the product rights, Amgen will pay royalties to BeOne on sales in China for a specified period. For product sales outside China, Amgen also pays royalties to BeOne.

For financial information about our significant collaborative arrangements, see Part IV—Note 9, Collaborations, to the Consolidated Financial Statements.

¹ We are also in a collaboration with AstraZeneca for the development of AMG 104. See Research and Development and Selected Product Candidates section above.

Human Capital Resources

Overview

Amgen's approach to human capital resource management starts with our mission to serve patients. We strive to serve patients by transforming the promise of science and biotechnology into therapies that have the power to restore health or save lives. The complexities of our industry along with the challenges of running an enterprise focused on the discovery, development, manufacture and commercialization of innovative medicines, require a highly engaged and committed workforce.

As of December 31, 2025, Amgen had approximately 31,500 staff members in over 50 countries, including approximately 13,000 staff members outside the United States, and we continue to experience overall low turnover rates. We also supplement our workforce with independent contractors, contingent workers and temporary workers, as needed. Outside of the United States, some of our employees are represented by unions or works councils. We consider our staff relations to be good, supported by regular assessments of staff engagement surveys on a wide range of topics (including flexible work environments, career development, and maintaining a culture of compliance). Our engagement scores were above general market benchmarks in 2025. We discuss the results of these surveys with our workforce and our Board of Directors. Reflecting our staff members' desire to retain a flexible approach to work, we offer a flexible workspace initiative that enables many employees to work together with their manager to determine the location that best enables their work at hand, supporting virtual work as well as working in person.

Compensation, Benefits and Development

Our approach to employee compensation and benefits is designed to deliver cash, equity and benefit programs that are competitive with those offered by leading companies in the biotechnology and pharmaceutical industries, and to attract, motivate and retain talent with a focus on encouraging performance, promoting accountability and adherence to our values and alignment with the interests of the Company's stockholders.

Our base pay program aims to compensate staff members relative to the value of the contributions of their role, which takes into account the skills, knowledge and abilities required to perform each position, as well as the experience brought to the job. We also provide annual incentive programs to reward our staff in alignment with achievement of Company-wide goals that are established annually and designed to drive aspects of our strategic priorities that support and advance our strategy across our Company and are intended to positively position us for both near- and long-term success. The majority of our staff members are also eligible for equity award grants under our long-term incentive program that are designed to align the interests of our staff members with those of our stockholders. For senior level staff, a significant proportion of equity award value is dependent on Company performance.

All staff participate in a regular performance measurement process through which staff receive performance and development feedback, and pay is aligned to performance. Our values and leadership attributes are an integral part of the performance assessments of our staff members, and these evaluations serve as an important information tool and basis for promotion and compensation decisions.

Our staff receive, and are guided by, regular trainings that help them understand what is expected of them. Further, to support the development of our staff, we provide a variety of programs, including leadership development programs, classroom-based and virtual instructor-led courses, and self-paced learning options as well as mentoring, networking and coaching opportunities. In 2025, we invested in general upskilling of our staff on skills for the future, including topics such as AI. We also maintain a global talent marketplace that enables staff to contribute to work beyond their core responsibilities, providing opportunities to broaden professional networks, support organizational priorities and develop additional skills.

Our benefit programs are generally broad-based, promote health and overall well-being and emphasize saving for retirement. All regular U.S. staff members are eligible to participate in the same core health and welfare and retirement savings plans. Other U.S. employee benefits include adoption assistance, paid parental leave programs, access to childcare, employee assistance programs, employee stock purchase plan, flexible spending accounts, life insurance, long-term care and business travel accident insurance, short and long-term disability benefits, wellness benefits and work-life resources and referrals. Comparable programs and benefits are available globally, with the same health and well-being intent, and consistent with local statutory requirements.

Our Compensation and Management Development Committee provides oversight of our compensation plans, policies and programs.

Total Workforce Health

Creating a safe and healthy workplace for our staff is an important priority at Amgen. Our goal is to have a world class safety record through safety leadership, engaged staff, risk management practices and integrating safety throughout our

business processes. We provide job-specific safety training tailored to each role, and to foster our safety culture, we implement a comprehensive safety program and reinforce desired safety behaviors, driving to understand and mitigate the root cause of safety incidents and manage and control variability. We track injuries and near-miss incidents through our incident tracking system, and we use leading indicators to assess the effectiveness of our safety programs and make course corrections as needed. Additionally, we perform formal executive management review of functional safety performance for Operations, Global Commercial Operations and R&D on a quarterly basis with a focus on identifying early signals and taking action to drive continuous improvement.

Our CRCC provides general oversight of our safety programs and initiatives.

Culture

We believe that our culture helps attract and retain a strong and engaged workforce informed by the varied experiences represented, which fosters innovation, collaboration and productivity as we execute on our mission to serve patients. A cross-functional, executive-level council, which includes our CEO, regularly discusses matters related to workplace culture and, as appropriate, takes actions to support and evolve Amgen's culture.

In our effort to attract and retain the best talent, we seek out and support talent across the globe. As part of our multidimensional hiring and talent development strategy, our Apprenticeship Program launched in 2023 in our Manufacturing and ATMOS functions and expanded to include another cohort of apprentices at our North Carolina site in January 2025. Our Apprenticeship Program is a skills-based approach and is designed to invest in our future workforce through attracting, hiring and upskilling non-four-year degreed talent in the United States. Through the Apprenticeship Program, we provide individuals with classroom-based and on-the-job training as well as mentorship opportunities needed to develop proficiency in targeted business areas and roles. Multiple cohorts are underway across various sites, including a new 18 month manufacturing apprenticeship at Amgen North Carolina, which serves as a key talent pipeline. We believe that our Apprenticeship Program and other skills-based approaches to hiring provide us with access to a larger pool of highly motivated and productive talent while also providing greater access to jobs in innovative sectors of the economy.

Our Compensation and Management Development Committee oversees our labor and employment policies, programs and initiatives, including those relating to our talent strategy and culture.

Information about Our Executive Officers

The executive officers of the Company as of February 13, 2026, are set forth below.

Mr. Robert A. Bradway, age 63, has served as a director of the Company since 2011 and Chairman of the Board of Directors since 2013. Mr. Bradway has been the Company's President since 2010 and Chief Executive Officer since 2012. From 2010 to 2012, Mr. Bradway served as the Company's President and Chief Operating Officer. Mr. Bradway joined the Company in 2006 as Vice President, Operations Strategy, and served as Executive Vice President and Chief Financial Officer from 2007 to 2010. Prior to joining the Company, Mr. Bradway was a Managing Director at Morgan Stanley in London, where, beginning in 2001, he had responsibility for the firm's banking department and corporate finance activities in Europe. Mr. Bradway has been a director of The Boeing Company, an aerospace company and manufacturer of commercial airplanes, defense, space and securities systems, since 2016. He has served on the board of trustees of the University of Southern California since 2014.

Dr. James E. Bradner, age 53, became Executive Vice President, Research and Development, in 2023 and also served as the Company's Chief Scientific Officer from 2023 to 2024. Prior to joining the Company, in 2023, Dr. Bradner was a clinician at the Dana-Farber Cancer Institute, a comprehensive cancer treatment and research institution, and a principal teaching affiliate at Harvard Medical School. From 2016 to 2022, Dr. Bradner served as President of the Novartis Institutes for BioMedical Research, the research and early development organizational unit of Novartis AG, where he was a member of the Executive Committee. Dr. Bradner previously served on the faculty at Harvard Medical School.

Mr. Murdo Gordon, age 59, became Executive Vice President, Global Commercial Operations, in 2018. Prior to joining the Company, Mr. Gordon was Chief Commercial Officer at Bristol-Myers Squibb Company (BMS), a pharmaceutical company, from 2016 to 2018. Mr. Gordon served as Head of Worldwide Markets at BMS from 2015 to 2016. Prior to this, Mr. Gordon served in a variety of leadership roles at BMS for more than 25 years.

Mr. Jonathan P. Graham, age 65, became Executive Vice President and General Counsel and Secretary in 2019. Mr. Graham joined the Company in 2015 to serve as Senior Vice President, General Counsel and Secretary. Prior to joining Amgen, from 2006 to 2015, Mr. Graham was Senior Vice President and General Counsel at Danaher Corporation. From 2004 to 2006, Mr. Graham was Vice President, Litigation and Legal Policy, at General Electric Company (GE). Prior to GE, Mr. Graham was a partner at Williams & Connolly LLP.

Mr. Peter H. Griffith, age 67, became Executive Vice President and Chief Financial Officer in 2020. Mr. Griffith joined the Company in 2019 as Executive Vice President, Finance. Prior to joining Amgen, Mr. Griffith was President of Sherwood Canyon Group, LLC, a private equity firm. From 1997 to 2019, Mr. Griffith was a partner at EY, an accounting and professional services firm, and served in a variety of senior leadership roles, with his last position being Global Vice Chair, Corporate Development. Prior to EY, Mr. Griffith was a Managing Director and head of the investment banking division of Wedbush Securities Inc.

Ms. Nancy A. Grygiel, age 58, became Senior Vice President and Chief Compliance Officer in 2020. Ms. Grygiel joined the Company in 2015 and has held leadership roles within the worldwide compliance and business ethics organization, including as Vice President, Compliance, from 2016 to 2020. Prior to joining Amgen, from 2011 to 2015, Ms. Grygiel served as Vice President, Compliance, Corporate & International, at Allergan, Inc. (Allergan). Prior to Allergan, Ms. Grygiel held several management positions at Mylan Pharmaceuticals, Inc.

Ms. Rachna Khosla, age 53, became Senior Vice President, Business Development, in 2021. Ms. Khosla joined the Company in 2013 and has held leadership roles in corporate development, licensing, and mergers and acquisitions, including as Vice President Business Development, from 2018 to 2021. Prior to joining the Company, Ms. Khosla was a Director at Lazard Ltd. (Lazard) responsible for healthcare mergers and acquisitions. Prior to Lazard, Ms. Khosla held various roles in investment banking (mergers and acquisitions) and corporate venture capital at Credit Suisse Group AG, Sanofi Aventis, Aventis Capital, J.P. Morgan Chase & Co., and Salomon Brothers, Inc.

Mr. Derek Miller, age 53, became Senior Vice President, Human Resources, in 2022. Mr. Miller joined the Company in 2003 and has held human resources leadership roles of increasing responsibility during his tenure. Mr. Miller's experience spans each of the Company's major business functions, including Vice President, Global Total Rewards, from 2020 to 2022 and Vice President, Human Resources, from 2018 to 2020.

Dr. David M. Reese, age 63, became the Company's inaugural Executive Vice President and Chief Technology Officer in 2023, responsible for accelerating the use of technology and artificial intelligence across the organization. From 2018 to 2023, Dr. Reese served as Executive Vice President, Research and Development. Dr. Reese joined the Company in 2005 and has held leadership roles in development, translational, and medical sciences, and discovery research, including as Senior Vice President, Translational Sciences and Oncology, from 2017 to 2018. Prior to joining Amgen, Dr. Reese was a cofounder, president, and chief medical officer of Translational Oncology Research International, a not-for-profit academic clinical research organization, and director of Clinical Research at the Breast Cancer International Research Group. Dr. Reese previously served on the faculty at the University of California, Los Angeles and the University of California, San Francisco.

Mr. Esteban Santos, age 58, became Executive Vice President, Operations, in 2016. Mr. Santos joined the Company in 2007 and has held a number of leadership roles at the Company in engineering, manufacturing, site operations and drug product, including Senior Vice President, Manufacturing, from 2013 to 2016. Prior to joining the Company, Mr. Santos served as Site General Manager of Johnson & Johnson's (J&J) Cordis operation in Puerto Rico. Prior to J&J, Mr. Santos held several management positions in GE's industrial and transportation businesses.

Geographic Area Financial Information

For financial information concerning the geographic areas in which we operate, see Part IV—Note 3, Revenues, and Note 12, Property, plant and equipment, to the Consolidated Financial Statements.

Investor Information

Financial and other information about us is available on our website at www.amgen.com. We make available on our website, free of charge, copies of our Annual Report on Form 10-K, Quarterly Reports on Form 10-Q, current reports on Form 8-K and amendments to those reports filed or furnished pursuant to Section 13(a) or 15(d) of the Exchange Act as soon as reasonably practicable after we electronically file such material with or furnish it to the U.S. Securities and Exchange Commission (SEC). In addition, we have previously filed registration statements and other documents with the SEC. Any document we file may be inspected without charge at the SEC's website at www.sec.gov. (These website addresses are not intended to function as hyperlinks, and the information contained in our website and in the SEC's website is not intended to be a part of this filing.)

Item 1A. RISK FACTORS

This report and other documents we file with the SEC contain forward-looking statements that are based on current expectations, estimates, forecasts and projections about us, our future performance, our business, our beliefs and our management's assumptions. These statements are not guarantees of future performance and involve certain risks, uncertainties and assumptions that are difficult to predict. You should carefully consider the risks and uncertainties our business faces. The risks described below are not the only ones we face. Our business is also subject to the risks that affect many other companies, such as employment relations, general economic conditions, geopolitical events and international operations. Further, additional risks not currently known to us or that we currently believe are immaterial may in the future materially and adversely affect our business, operations, liquidity and stock price.

SUMMARY

Risks Related to Government Regulations and Third-Party Policies

- Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability.
- Guidelines and recommendations published by various organizations can reduce the use of our products.
- We could be subject to additional tax liabilities, including from an adverse outcome in our ongoing tax dispute with the IRS and other tax examinations, enactment of the OECD minimum corporate tax rate agreement and the adoption and interpretation of new tax legislation, including OB3. Such tax liabilities could adversely affect our profitability and results of operations.
- Our business may be affected by litigation and government investigations.

Risks Related to Economic Conditions and Operating a Global Business

- Our efforts to collaborate with or acquire other companies, products, or technology, and to integrate the operations of companies or to support the products or technology we have acquired, may not be successful, and may result in unanticipated costs, delays or failures to realize the benefits of the transactions.
- A breakdown of our information technology systems, cyberattack or information security breach could significantly compromise the confidentiality, integrity and availability of our information technology systems, network-connected control systems and/or our data, interrupt the operation of our business and/or affect our reputation.
- Our sales and operations are subject to the risks of doing business internationally, including in new or emerging markets.
- We may not be able to access the capital and credit markets on terms that are favorable to us, or at all.

Risks Related to Competition

- Our products face substantial competition and our product candidates are also likely to face substantial competition.
- Our intellectual property positions may be challenged, invalidated or circumvented, or we may fail to prevail in current and future intellectual property litigation.
- We currently face competition from biosimilars and generics and expect to face increasing competition from biosimilars and generics in the future.
- Concentration of sales at certain of our wholesaler distributors, and consolidation of private payers, such as insurers, and PBMs has negatively affected, and may continue to negatively affect, our business.

Risks Related to Research and Development

- We may not be able to develop commercial products despite significant investments in R&D.
- We must conduct clinical trials in humans before we commercialize and sell any of our product candidates or existing products for new indications.
- Our current products and products in development cannot be sold without regulatory approval.

- Some of our products are used with drug delivery or companion diagnostic devices that have their own regulatory, manufacturing and other risks.
- Some of our pharmaceutical pipeline and our commercial product sales rely on collaborations with third parties, which may adversely affect the development and sales of our products.

Risks Related to Operations

- We perform a substantial majority of our commercial manufacturing activities at our facility in the U.S. territory of Puerto Rico and a substantial majority of our clinical manufacturing activities at our facility in Thousand Oaks, California; significant disruptions or production failures at these facilities could significantly impair our ability to supply our products or continue our clinical trials.
- We rely on third-party suppliers for certain of our raw materials, medical devices and components.
- Manufacturing difficulties, disruptions or delays could limit supply of our products and limit our product sales.
- Our business and operations may be negatively affected by the failure, or perceived failure, of achieving our sustainability objectives.
- The effects of global climate change and related natural disasters could negatively affect our business and operations.

General Risk Factors

- Global economic conditions may negatively affect us and may magnify certain risks that affect our business.
- Our stock price is volatile.

RISKS RELATED TO GOVERNMENT REGULATIONS AND THIRD-PARTY POLICIES

Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability.

Sales of our products depend on the availability and extent of coverage and reimbursement from third-party payers, including government healthcare programs and private insurance plans. Governments and private payers continue to pursue initiatives to manage drug utilization and contain costs. Payers are increasingly focused on costs, which has resulted, and is expected to continue to result, in lower reimbursement rates for our products and/or narrower patient populations for which payers will reimburse. Continued intense public scrutiny of the price of drugs and other healthcare costs, together with payer dynamics, have limited, and are likely to continue to limit, our ability to set or adjust the price of our products based on their value, which can have a material adverse effect on our business. In the United States, a number of legislative and regulatory proposals have been introduced and/or signed into law to lower drug prices. These include the IRA that enables the U.S. government to set prices for certain drugs in Medicare, redesigns Medicare Part D benefits to shift a greater proportion of the costs to manufacturers and health plans, and enables the U.S. government to impose penalties if drug prices are increased at a rate faster than inflation (IRA Inflation Penalties). On July 4, 2025, OB3 was enacted and included several changes to Medicare, Medicaid and Affordable Care Act policies, including provisions affecting eligibility, that, when implemented, are expected to adversely affect coverage and reimbursement for our products. On May 12, 2025, the Administration issued the Most-Favored-Nations (MFN) Prescription Drug Pricing Executive Order (MFN EO) aimed at using price benchmarks from other developed countries to set U.S. pricing targets. Subsequently, on July 31, 2025 the Administration sent letters to many pharmaceutical manufacturers, including Amgen (the July MFN Letter) as further described below, outlining steps that such manufacturers could take to advance actions consistent with elements of the MFN EO. In December 2025, we announced that we are taking actions that satisfy the components outlined in the July MFN Letter, including the Administration's MFN pricing requests. Further, the Administration has called on Congress to enact legislation that would codify the terms that the Administration arrived at with recipients of the July MFN Letter (the MFN Terms). The details of such legislative framework are unknown and, if enacted, such legislation could apply to a broader range of products, payers or pricing arrangements for a longer period than those resulting from the MFN Terms. Additional proposals focused on drug pricing continue to be debated, and additional executive orders or regulatory initiatives focused on drug pricing and competition may be adopted and implemented in some form. It remains unclear what further policies, legislation and/or actions the Administration, Congress, or state governments will advance with respect to other drug pricing proposals or other healthcare regulations affecting pharmaceuticals, including the MFN EO, IRA and OB3 implementation, trade policies, or state laws affecting the 340B Program or Medicaid reimbursement that could ultimately be adopted more broadly. To the extent such actions reduce or modify coverage or reimbursement for our products, increase rebates or other costs, constrain pricing decisions, or otherwise limit product use, they would have an adverse effect on our business and results of operations.

—Changing U.S. federal coverage and reimbursement policies and practices have affected, and are likely to continue to affect, access to, pricing of, and sales of our products

A substantial proportion of our U.S. business relies on reimbursement from federal government healthcare programs and commercial insurance plans regulated by federal and state governments. See Item 1. Business—Reimbursement. Our business has been, and will continue to be, affected by legislative actions changing U.S. federal reimbursement policy. For example, the IRA includes provisions requiring mandatory pricing in Medicare for certain drugs under Parts B and D (starting with 10 drugs effective January 1, 2026, adding 15 in 2027 and 2028, and adding 20 in 2029 and subsequent years such that, by 2031, approximately 100 drugs would be subject to such set prices). CMS has set Medicare Part D prices for ENBREL, effective January 1, 2026, and Otezla, effective January 2027, in each case at significantly lower prices that are expected to negatively impact their profitability in Medicare. See Part II, Item 7. Management’s Discussion and Analysis of Financial Condition and Results of Operations—Results of operations—*Product sales*. Further, CMS has issued guidance that allows for the re-setting of prices for drugs for which it previously set a price. Depending on the growth and success of our medicines, other of our medicines may also be subject to selection by CMS in the next, or in a future, cycle of mandatory Medicare price setting, we may be required to accept a price set by the government for Medicare using the process that was applied to ENBREL and Otezla. On April 15, 2025, the Administration issued an executive order (the April 2025 EO) that, among other directives, directs HHS to work with Congress to align the treatment of small molecule drugs and biologics in the Medicare price setting program under the IRA. It is currently unclear how such modifications would affect the timeframe in which Medicare price setting becomes applicable for selected drugs or biologics. Also under the IRA, Medicare Part D was redesigned to cap beneficiary out-of-pocket costs and reduce Federal reinsurance in the catastrophic phase; increasing cost-sharing obligations for Part D plans and manufacturers, including by requiring manufacturer discounts. Further, the IRA inflation penalties allow CMS to collect rebates from manufacturers if Medicare price increases outpace inflation, and several of our products have been subject to such IRA inflation rebates. The IRA’s Medicare price setting and Medicare redesign have had, and are likely to have, an adverse effect on our sales, our business and our results of operations, and such impact is expected to increase through the end of the decade and will depend on factors including the extent of our portfolio’s exposure to Medicare reimbursement, the rate of inflation over time, the number of our products selected for Medicare price setting and the timing of market entry of generic or biosimilar competition. Further, following the enactment of the IRA, the environment remains dynamic, and U.S. policymakers continue to demonstrate interest in health care and drug pricing changes as well as potential changes affecting intellectual property. For example, in April 2024, CMS finalized policy changes that will give Part D plans more flexibility to substitute biosimilars for innovator products on formularies in 2025. Implementation of OB3 also may impact access to and reimbursement of our products. For example, the Congressional Budget Office has projected that the OB3 will result in significant reductions in federal Medicaid spending over the next decade and an increase in the number of people without health insurance. These developments would place greater stress on state budgets and hospital finances, and could result in reduced access to medicines, additional pressure to further discount medicines and further growth of 340B Program utilization. The MFN EO directs HHS to pursue pricing policies that align U.S. drug prices with the prices available in certain comparably developed countries and directs a range of actions to advance that objective, including through regulatory, trade and other policy measures. In July 2025, the Administration delivered the July MFN Letter to us and a number of other manufacturers that called for drug manufacturers to: 1) extend MFN pricing to Medicaid; 2) guarantee MFN pricing to Medicaid, Medicare and commercial payers on all newly launched drugs; 3) use future increased revenues from outside the U.S. to lower U.S. drug prices; and 4) participate in direct-to-patient models to provide MFN pricing for certain drugs. Since the issuance of the July MFN Letter, most recipients have announced that they have reached agreement with the Administration to address the matters described in the letter. In December 2025, we announced that we are taking actions that satisfy the components outlined in the July MFN Letter, including the Administration’s MFN pricing requests. We also announced the expansion of our direct-to-patient program. While this development reflects ongoing engagement on pricing policy, the ultimate effects on our pricing, reimbursement, net sales and profitability remain uncertain in light of evolving regulatory and policy expectations.

We also face risks related to the reporting of pricing data that affects reimbursement of and discounts provided for our products. U.S. government price reporting regulations are complex and may require biopharmaceutical manufacturers to update certain previously submitted data. If our submitted pricing data are incorrect, we may become subject to substantial fines and penalties or other government enforcement actions, which could have a material adverse effect on our business and results of operations. In addition, as a result of restating previously reported price data, we may be required to pay additional rebates and provide additional discounts.

—Changing reimbursement and pricing actions in various states have negatively affected, and may continue to negatively affect, access to, and have affected, and may continue to affect, sales of our products

At the state level, legislation, government actions and ballot initiatives can also affect how our products are covered and reimbursed and/or create additional pressure on our pricing decisions. Existing and proposed state pricing laws, which may move forward more rapidly than similar efforts at the federal level, have added complexity to the pricing of drugs. A number of states have adopted, and many other states are considering, PDABs, drug importation programs, reference pricing schemes and

other drug pricing actions, including proposals designed to require biopharmaceutical manufacturers to report to the state proprietary pricing information or provide advance notice of certain price increases.

States continue to pursue laws related to price controls, referencing the IRA and seeking to regulate and prohibit restrictions on the 340B Program. For example, following the passage of the IRA, bills have been proposed in multiple states that would apply the drug price caps set by HHS for Medicare to drug prices in an individual state, and such references to IRA price caps have also been included in PDAB legislation. For Medicaid patients, states have established a Medicaid drug spending cap (New York) and implemented a new review and supplemental rebate negotiation process (Massachusetts). Eight states (Colorado, Maine, New Hampshire, New Jersey, Maryland, Minnesota, Oregon and Washington) have enacted laws that establish PDABs to identify drugs that pose affordability challenges, and four such states include authority for the state PDABs to set upper payment limits on certain drugs for in-state patients, payers and providers. In 2025, Maryland expanded the scope of its PDAB law to include the commercial market. The eight states with enacted PDAB laws are in various phases of implementation, with Colorado's PDAB being the furthest along. The Colorado PDAB deemed three of five drugs "unaffordable," including ENBREL, and in October 2025 the Colorado PDAB established an Upper Payment Limit (UPL) substantially lower than the wholesale acquisition cost of ENBREL that would be generally applicable to all formulations of ENBREL, effective no earlier than January 1, 2027, and will be reviewed annually. On July 16, 2025, Washington state's PDAB selected ENBREL for one of its first affordability reviews. Following the timeline and process established by the state for such affordability review, the manufacturer and the PDAB will undertake a number of required interactions. However, the Washington state PDAB may not establish a UPL for any prescription drug before January 1, 2027. Further, inappropriate expanded utilization of the 340B Program from broadened application of the 340B discounts has had, and is expected to continue to have, a negative impact on the Company's product sales, business and results of operations. Twenty states (Louisiana, Arkansas, West Virginia, Minnesota, Mississippi, Missouri, Maryland, North Dakota, South Dakota, Utah, Nebraska, New Mexico, Colorado, Tennessee, Oregon, Vermont, Hawaii, Oklahoma, Rhode Island and Maine) have enacted laws with mandates on manufacturers participating in the 340B Program, and, in 2025, no fewer than 30 states have introduced similar legislation. These bills vary, but typically include provisions restricting a manufacturer's ability to direct drugs in 340B channels, recognizing 340B contract pharmacies and a prohibition on requiring the inclusion of 340B claims modifiers. With OB3's reductions to federal Medicaid funding to states, increased pressure is anticipated for providers to find and preserve existing revenue sources at the state level, which may result in increased use of 340B contract pharmacy mandates. In *Genesis Health Care, Inc. v. Becerra*, the U.S. District Court for the District of South Carolina issued an order in November 2023 enjoining the Health Resources and Services Administration from enforcing a more restrictive interpretation against Genesis Health Care as to who qualifies as a patient under the 340B Program, which could, if adopted more broadly, affect the scope of eligibility for 340B discounts. Since this decision, various courts have reached differing conclusions on challenges to state laws regulating aspects of the 340B Program, with some courts declining to enjoin such laws and others granting relief to challengers. Certain appellate courts have issued decisions both upholding certain state 340B statutes and affirming the denial of preliminary injunctive relief to manufacturers, while litigation and appeals concerning the validity, interpretation, and enforcement of these laws remain ongoing.

Additionally, in 2024, the FDA authorized Florida to move forward with its importation program proposal, though the state has not yet completed any significant steps towards importation within the two-year authorization window. Colorado, Maine, New Hampshire, New Mexico, Texas and Vermont have also enacted state importation laws, and some have submitted plans for approval to the FDA. Other states could adopt similar approaches or could pursue different policy changes in a continuing effort to reduce their costs. Further, the April 2025 EO also directs HHS to, within 90 days, streamline and improve the drug importation program to ease the process for states to obtain drug importation approvals. On May 21, 2025, the FDA issued a press release indicating it was taking steps to enhance state importation programs and would offer individual states and tribes the opportunity to submit draft proposals for pre-review and to meet with the agency to obtain initial feedback prior to formally submitting importation proposals. While under federal law biologics remain exempt from such state importation activities, our small molecule products could be impacted by these initiatives.

Ultimately, existing or future state government actions or ballot initiatives may also have a material adverse effect on our product sales, business and results of operations.

—*U.S. commercial payer actions have affected, and may continue to affect, access to and sales of our products*

Payers, including healthcare insurers, PBMs, integrated healthcare delivery systems (vertically-integrated organizations built from consolidations of healthcare insurers and PBMs) and group purchasing organizations, are continuing to seek ways to further reduce their costs. With increasing frequency, payers are adopting benefit plan changes that shift a greater proportion of drug costs to patients. Such measures include more limited benefit plan designs, high deductible plans, higher patient co-pay or coinsurance obligations and more significant limitations on patients' use of manufacturer commercial co-pay assistance programs. Further, government regulation of payers may affect these trends. Payers, including PBMs, have sought, and continue to seek, price discounts or rebates in connection with the placement of our products on their formularies or those they

manage, and to also impose restrictions on access to, or usage of, our products (such as Step Therapy), require that patients receive the payer's prior authorization before covering the product, and/or to exclude certain indications for which our products are approved.

Further, significant consolidation in the health insurance industry has resulted in a few large insurers and PBMs, which places greater pressure on pricing and usage negotiations with biopharmaceutical manufacturers, significantly increasing discount and rebate requirements and limiting patient access and usage. See —*Concentration of sales at certain of our wholesaler distributors, and consolidation of private payers, such as insurers, and PBMs has negatively affected, and may continue to negatively affect, our business.* This high degree of consolidation among insurers, PBMs and other payers, including integrated healthcare delivery systems and/or with specialty or mail-order pharmacies and pharmacy retailers, has increased the negotiating leverage such entities have over us and other biopharmaceutical manufacturers and has resulted in greater price discounts, rebates and service fees realized by those payers from our business. Our business is also affected by policies implemented by private healthcare entities that process Medicare claims, including Medicare Administrative Contractors. Each of CVS, Express Scripts and United Health Group (among the top six integrated health plans and PBMs) have Rebate Management Organizations that further increase their leverage to negotiate deeper discounts on their behalf and for the benefit of their other customers. Ultimately, additional discounts, rebates, fees, coverage changes, plan changes, restrictions or exclusions imposed by these commercial payers could have a material adverse effect on our product sales, business and results of operations. See —*Concentration of sales at certain of our wholesaler distributors, and consolidation of private payers, such as insurers, and PBMs has negatively affected, and may continue to negatively affect, our business.*

—*Government and commercial payer actions outside the United States have affected and will continue to affect access to and sales of our products*

Outside the United States, we expect countries will also continue to take actions to reduce their drug expenditures and to reduce intellectual property protections. See Item 1. Business—Reimbursement. Pressures to decrease drug expenditures may intensify as governments take actions to address budgets strained by high inflation and weak economic conditions, including in Europe, where sustained fiscal pressures continue to challenge public healthcare systems. In addition, policies under consideration or adopted in the United States that reference or tie drug prices to those paid in foreign jurisdictions, including through most-favored-nation or similar pricing approaches, could increase the significance of pricing decisions in such foreign jurisdictions. Further, the EU is currently undergoing a review and revision of its general pharmaceutical legislation that, while full implementation is not expected before 2027, has led to proposals that would reduce intellectual property protection for new products (including potentially shortening the duration of regulatory data exclusivity and orphan drug exclusivity protections), as well as change the reimbursement and regulatory landscape. International reference pricing has been widely used by many countries outside the United States to control costs. International reference pricing policies can change quickly and frequently and may not reflect differences in the burden of disease, indications, market structures or affordability across countries or regions. Other expenditure control practices, including the use of revenue clawbacks, rebates and caps on product sales, are also used in various foreign jurisdictions. In addition, countries may refuse to reimburse, or may restrict the reimbursed population for a product, when their national health technology assessments do not consider a medicine to demonstrate sufficient clinical benefit beyond existing therapies or to meet certain cost effectiveness thresholds. For example, despite the EMA's approval of Repatha for the treatment of patients with established atherosclerotic disease, prior to 2020, the reimbursement of Repatha in France was limited to a narrower patient population (such as those with homozygous familial hypercholesterolemia (HoFH)) following a national health technology assessment. Many countries decide on reimbursement between potentially competing products through national or regional tenders that often result in one product receiving most, or all of, the sales in that country or region. Failure to obtain coverage and reimbursement for our products, a deterioration in their existing coverage and reimbursement, or a decline in the timeliness or certainty of payment by payers to hospitals and other providers, has negatively affected, and may further negatively affect, the ability or willingness of healthcare providers to prescribe our products for their patients and otherwise negatively affect the use of our products or the prices we realize for them. Such failures and changes have had, and could in the future have, a material adverse effect on our product sales, business and results of operations.

Guidelines and recommendations published by various organizations can reduce the use of our products.

Government agencies promulgate regulations and guidelines directly applicable to us and to our products. Professional societies, practice management groups, insurance carriers, physicians' groups, private health and science foundations and organizations involved in various diseases also publish guidelines and recommendations to healthcare providers, administrators and payers, as well as patient communities. Recommendations by government agencies or other groups and organizations may relate to such matters as usage, dosage, route of administration and use of related therapies. In addition, a growing number of organizations are providing assessments of the value and pricing of biopharmaceutical products, and even organizations whose guidelines have historically been focused on clinical matters have begun to incorporate analyses of the cost effectiveness of various treatments into their treatment guidelines and recommendations. Value assessments may come from private organizations that publish their findings and offer recommendations relating to the products' reimbursement by government and

private payers. Some companies and payers have announced pricing and payment decisions based in part on the assessments of private organizations. In addition, government health technology assessment organizations in many countries make reimbursement recommendations to payers in their jurisdictions based on the clinical effectiveness, cost-effectiveness and service effects of new, emerging and existing medicines and treatments. Such health technology assessment organizations have *recommended*, and may in the future recommend, reimbursement for certain of our products for a narrower indication than was approved by applicable regulatory agencies or may recommend against reimbursement entirely. See *Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability*. The EU has adopted and begun implementing Regulation 2021/2282 on Health Technology Assessment, which establishes a permanent framework for cooperation among EU member states on joint clinical assessments and scientific consultations of health technologies, with key provisions that became applicable in January 2025, and further phased implementation continuing through 2030. These and other such recommendations or guidelines may affect our reputation, and any recommendations or guidelines that result in decreased use, dosage or reimbursement of our products could have a material adverse effect on our product sales, business and results of operations. In addition, the perception by the investment community or stockholders that such recommendations or guidelines will result in decreased use and dosage of our products could adversely affect the market price of our common stock.

We could be subject to additional tax liabilities, including from an adverse outcome in our ongoing tax dispute with the IRS and other tax examinations, enactment of the OECD minimum corporate tax rate agreement and the adoption and interpretation of new tax legislation, including OB3. Such tax liabilities could adversely affect our profitability and results of operations.

We are subject to income and other taxes in the United States and other jurisdictions in which we do business. As a result, our provision for income taxes is derived from a combination of applicable tax rates in the various places we operate. Significant judgment is required for determining our provision for income tax.

One or more of our legal entities file income tax returns in the U.S. federal jurisdiction, various U.S. state jurisdictions and foreign jurisdictions. Our income tax returns are routinely examined by tax authorities in those jurisdictions. Significant disputes can and have arisen with tax authorities involving issues regarding the timing and amount of deductions, the use of tax credits and allocations of income and expenses among various tax jurisdictions because of differing interpretations of tax laws, regulations and relevant facts, and such tax authorities (including the IRS) are becoming more aggressive in its audits and are particularly focused on such matters.

In 2017, we received an RAR and a modified RAR from the IRS for the years 2010–2012, proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico. We disagreed with the proposed adjustments and calculations, and in 2021, we filed a petition in the U.S. Tax Court to contest two duplicate Statutory Notices of Deficiency (Notices) for the years 2010–2012. The Notices seek to increase our U.S. taxable income for the years 2010–2012.

In 2020, we received an RAR and a modified RAR from the IRS for the years 2013–2015, also proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico similar to those proposed for the years 2010–2012. We disagreed with the proposed adjustments and calculations, and in 2022, we filed a petition in the U.S. Tax Court to contest a Notice for the years 2013–2015. The Notice seeks to increase our U.S. taxable income for the years 2013–2015 and asserts penalties.

We firmly believe that the IRS positions set forth in the 2010–2012 and 2013–2015 Notices are without merit. We continue to contest the 2010–2012 and 2013–2015 Notices through the judicial process. The two cases were consolidated in 2022. The trial began on November 4, 2024 and concluded on January 17, 2025. The parties filed opening post-trial briefs on June 13, 2025, and the Court held oral argument on July 16, 2025. The parties filed post-trial reply briefs on October 10, 2025. The Company expects a decision from the U.S. Tax Court no earlier than the second half of 2026.

We are currently also under examination by the IRS for the years 2016–2018 with respect to issues similar to those for the 2010 through 2015 period. As a result of the audit, the IRS could propose adjustments or penalties that, in all or part, relate to issues in the 2010–2015 dispute. We expect that the IRS will begin its audit of 2019–2022 in the first half of 2026, and we believe that it may seek to continue to audit similar issues related to the allocation of income between the United States and our foreign jurisdictions. In addition, we are under examination by a number of state and foreign tax jurisdictions.

Final resolution of these complex tax matters is not likely within the next 12 months. We continue to believe our accrual for income tax liabilities is appropriate based on past experience, interpretations of tax law, application of the tax law to our facts and judgments about potential actions by tax authorities; however, due to the complexity of the provision for income taxes and uncertain resolution of these matters, the ultimate outcome of any tax matters may result in payments substantially greater than amounts accrued and could have a material adverse effect on the results of our operations. See Part II, Item 7.

Our provision for income taxes and results of operations in the future could be adversely affected by changes to our operating structure, changes in the mix of income and expenses in countries with differing tax rates, changes in the valuation of deferred tax assets and liabilities and changes in applicable tax laws, regulations or administrative interpretations thereof. The 2017 Tax Act is complex and a large volume of regulations and guidance has been issued and could be subject to different interpretations. We could face audit challenges to our application of the 2017 Tax Act.

As previously reported, the OECD reached an agreement to align countries on a minimum corporate tax rate and an expansion of the taxing rights of market countries. Select individual countries, including the United Kingdom, EU member countries, and Singapore, have enacted the global minimum tax agreement that took effect starting in 2024. On January 5, 2026, the OECD issued additional administrative guidance related to the global minimum tax agreement that exempts U.S. companies from extra territorial minimum taxes effective January 1, 2026. We are monitoring the potential 2026 impact of such administrative guidance as jurisdictions enact the new rules. Delays, modifications or repeal, in whole or in part, of the administrative guidance or the OECD agreement itself, either by all OECD participants or unilaterally by individual countries, could result in negative impacts on our overall tax rate.

OB3 includes significant provisions related to taxation, such as the permanent extension of certain expiring provisions of the 2017 Tax Act, and modifications to the international tax framework. This legislation has multiple effective dates, with certain provisions effective in 2026 and beyond. While we have incorporated the impact of OB3 in our 2025 financial statements, the legislation is complex and guidance could be issued that could impact our 2025 or future U.S. taxes.

Changes to existing tax law in the United States, the U.S. territory of Puerto Rico or other jurisdictions, including the changes and potential changes discussed above, could result in tax increases where we do business and could have a material adverse effect on the results of our operations.

Our business may be affected by litigation and government investigations.

We and certain of our subsidiaries are involved in legal proceedings. See Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements. Civil and criminal litigation is inherently unpredictable, and the outcome can result in costly verdicts, fines and penalties, exclusion from federal healthcare programs and/or injunctive relief that affect how we operate our business. Defense of litigation claims can be expensive, time consuming and distracting, and it is possible that we could incur judgments or enter into settlements of claims for monetary damages or change the way we operate our business, which could have a material adverse effect on our product sales, business and results of operations. In addition, product liability is a major risk in testing and marketing biotechnology and pharmaceutical products. We may face substantial product liability exposure in human clinical trials and for products we sell after regulatory approval. Product liability claims, regardless of their merits, could be costly and divert management's attention and could adversely affect our reputation and the demand for our products. We and certain of our subsidiaries have previously been, and currently are, named as defendants in product liability actions for certain of our products.

We are also involved in government investigations that arise in the ordinary course of our business. In recent years, there has been a trend of increasing government investigations and litigations against companies operating in our industry, both in the United States and around the world. See *Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability*. Our business activities outside of the United States are subject to the FCPA and similar antibribery or anticorruption laws, regulations or rules of other countries in which we operate, including the U.K. Bribery Act. We cannot ensure that all our employees, agents, contractors, vendors, licensees, partners or collaborators will comply with all applicable laws and regulations. We entered into a corporate integrity agreement with the DOJ and the Office of Inspector General of the HHS to settle certain allegations relating to our support of independent charitable organizations that provide patients with financial assistance to access their medicines that required us to maintain a corporate compliance program and to undertake a set of defined corporate integrity obligations that we completed in 2024. While we fully complied with all of our obligations under the corporate integrity agreement, we may be subject to future corporate integrity agreements and failure to comply could result in substantial penalties and potential exclusion from government healthcare programs. We may also see new government investigations of or actions against us citing novel theories of recovery. For example, prosecutors are placing greater scrutiny on patient support programs, including commercial copay assistance programs, and further enforcement actions and investigations regarding such programs could limit our ability to provide co-pay assistance to commercial patients. Greater scrutiny has also been placed on sponsorships, speaker programs and other arrangements where healthcare professionals receive remuneration, travel or other value to participate in certain events, and further enforcement actions could adversely affect our ability to participate in such arrangements. Any of these results could have a material adverse effect on our business and results of operations.

RISKS RELATED TO ECONOMIC CONDITIONS AND OPERATING A GLOBAL BUSINESS

Our efforts to collaborate with or acquire other companies, products, or technology, and to integrate the operations of companies or to support the products or technology we have acquired, may not be successful, and may result in unanticipated costs, delays or failures to realize the benefits of the transactions.

We seek innovation through significant investment in both internal R&D and external transactions, including collaborations, partnerships, alliances, licenses, joint ventures, mergers and acquisitions (collectively, acquisition activity). Acquisition activities may be subject to regulatory approvals or other requirements that are not within our control. Antitrust scrutiny by regulatory agencies and changes to regulatory approval process in the U.S. and foreign jurisdictions may cause approvals to take longer than anticipated to obtain, not be obtained at all, or contain burdensome conditions, which may jeopardize, delay or reduce the anticipated benefits of acquisitions to us and could impede the execution of our business strategy. There can be no assurance that such regulatory or other approvals will be obtained or that all closing conditions required in connection with our acquisition activities will be satisfied or waived, which could result in us being unable to complete the planned acquisition activities.

Acquisition activities are complex, time consuming and expensive and may result in unanticipated costs, delays, or other operational or financial problems related to integrating the acquired company and business with our company, which may divert our management's attention from other business issues and opportunities may prevent us from realizing the anticipated benefits of such transactions within the expected timeframe or at all.

We may pay substantial amounts of cash, incur debt or issue equity securities to pay for acquisition activities, which could adversely affect our liquidity or result in dilution to our stockholders, and could adversely affect our credit ratings and cost of capital. For example, the primary sources of funds for our acquisition of Horizon were those received from our \$24 billion of senior notes issued on March 2, 2023, together with the \$4 billion drawn down from our term loan facility, of which we repaid \$2.2 billion, and while the Company currently has investment grade credit ratings and substantially returned to its pre-acquisition capital structure, when the Company incurred this substantial additional indebtedness the credit rating agencies downgraded our credit ratings. Further, failures or difficulties in integrating or retaining key personnel, or in integrating the operations of the businesses, products or assets we acquire (including related technology, research, development and commercial operations, compliance programs, manufacturing, distribution and general business operations and procedures and sustainability activities) may adversely affect our ability to realize the benefits of the transaction or grow our business, and may result in asset impairment or restructuring charges. These and other challenges may arise in connection with our acquisitions, including our acquisitions of ChemoCentryx, Horizon and Dark Blue Therapeutics and/or our collaborations with BeOne and Kyowa Kirin, or with other acquisition activities, which could have a material adverse effect on our business, results of operations and stock price.

We may not realize the anticipated strategic benefits of our acquisitions. Our assumptions and estimates about the future revenue growth of acquired products may prove to be incorrect. Business integrations generally are complex, time consuming and expensive, and we may experience unanticipated costs, delays or other operational or financial challenges. These integration efforts may also divert our management's attention and resources away from other business operations, which may disrupt to some degree our ongoing business. Failure to successfully fully integrate acquired businesses into ours and/or achieve anticipated strategic benefits may result in our incurring significant asset impairment or restructuring charges, and could have a material adverse effect on our business, results of operations and stock price.

A breakdown of our information technology systems, cyberattack or information security breach could significantly compromise the confidentiality, integrity and availability of our information technology systems, network-connected control systems and/or our data, interrupt the operation of our business and/or affect our reputation.

To achieve our business objectives, we rely on sophisticated information technology systems, including hardware, software, technology infrastructure, online sites and networks for both internal and external operations, mobile applications, cloud services, artificial intelligence (AI)-enabled tools and systems, and network-connected control systems, some of which are managed, hosted, provided or serviced by third parties. Internal or external events that compromise the confidentiality, integrity and availability of our systems and data may significantly interrupt the operation of our business, result in significant costs and/or adversely affect our reputation.

Our information technology systems, including AI-enabled tools and systems are highly integrated into our business, including our R&D efforts, our clinical and commercial manufacturing processes and our product sales and distribution processes. Further, as the majority of our employees work remotely for some portion of their jobs in our hybrid work environment, our reliance on our and third-party information technology systems has increased substantially and is expected to continue to increase. Remote and hybrid working arrangements, including those of many third-party providers, can increase cybersecurity risks due to the challenges associated with managing remote computing assets and security vulnerabilities that are

present in many non-corporate and home networks. The complexity and interconnected nature of software, hardware and our systems make them vulnerable to breakdown or other service interruptions, and to software errors or defects, misconfiguration and other security vulnerabilities. For example, in July 2024, businesses worldwide were affected by an information technology outage due to a faulty software update issued by a cybersecurity firm. Although our systems and operations were temporarily affected by the outage, the impact of this firm's faulty update on the Company was immaterial to our business operations. However, there can be no assurance that a future similar incident would not result in a material adverse effect on our business or results of operations. Upgrades or changes to our systems or the software that we use have resulted and we expect, in the future, will result in the introduction of new cybersecurity vulnerabilities and risks. In the past, we have identified security vulnerabilities introduced into our information systems arising from flaws in third-party software that we had purchased and installed, which required us to apply emergency patches to certain systems. While we did not experience any significant adverse effects as a result of these vulnerabilities, there can be no assurance that we will timely identify and address future vulnerabilities. Our systems are also subject to frequent perimeter network reconnaissance and scanning, phishing and other cyberattacks. For example, as a result of our cybersecurity monitoring of the Horizon legacy information systems, we detected phishing activity in the accounts of two Horizon executives. These accounts were de-activated, the incidents were investigated and the determination was made separately by both our internal cybersecurity team and our external digital forensics and incident response supplier that no confidential information had been exfiltrated, and the incidents are now closed. As the cyber-threat landscape evolves, these attacks are growing in frequency, sophistication, and intensity, and are becoming increasingly difficult to detect and increasingly sophisticated in using techniques and tools, including AI, that circumvent security controls, evade detection and remove forensic evidence. Such attacks include the use of harmful and virulent malware, including ransomware or other denials of service, which can be deployed through various means, including the software supply chain, e-mail, malicious websites and/or the use of social engineering/phishing, and/or AI to orchestrate and automate sophisticated cyberattacks, including the documented instance in which an AI agent was used to conduct a large-scale intrusion campaign, potentially lowering the barriers to high-speed, high-volume attacks.

We have also experienced denial of service attacks against our network, and, although such attacks did not succeed, there can be no assurance that our efforts to guard against the wide and growing variety of potential attack techniques will be successful in the future. Attacks such as those experienced by government entities (including those that approve and/or regulate our products, such as the EMA) and other multi-national companies, including some of our peers, could leave us unable to utilize key business systems or access or protect important data, and could have a material adverse effect on our ability to operate our business, including developing, gaining regulatory approval for, manufacturing, selling and/or distributing our products. For example, in 2017, a pharmaceutical company experienced a cyberattack involving virulent malware that significantly disrupted its operations, including its research and sales operations and the production of some of its medicines and vaccines. As a result of the cyberattack, its orders and sales for certain products were negatively affected. In late 2020, SolarWinds Corporation, a leading provider of software for monitoring and managing information technology infrastructure, disclosed that it had suffered a cybersecurity incident whereby attackers had inserted malicious code into legitimate software updates for its products that were installed by myriad private and government customers, enabling the attackers to access a backdoor to such systems. We also rely on third-party providers for certain identity, authentication and access management services. In the past, a security incident affecting such a third-party provider exposed several hundred corporate customers to potential unauthorized access to systems and data. Although that breach did not have a significant effect on our business, there can be no assurance that a similar future breach would not result in a material adverse effect on our business or results of operations.

Our systems also contain and use a high volume of sensitive data, including intellectual property, trade secrets and other proprietary business information, financial information, regulatory information, strategic plans, sales trends and forecasts, litigation materials and/or personal identifiable information belonging to us, our staff, our patients, customers and/or other parties. In some cases, we use third-party service providers to collect, process, store, manage or transmit such data, which has increased our risk. Intentional or inadvertent data privacy or security breaches (including cyberattacks) resulting from attacks or lapses by employees, service providers (including providers of information technology-specific services), business partners, nation states (including groups associated with or supported by foreign intelligence agencies), organized crime organizations, "hacktivists" or others, create risks that our sensitive data may be exposed to unauthorized persons, our competitors or the public. Malicious actors, including those working under state-sponsored campaigns, have sought employment, often in remote information technology roles, as a means to gain inside access at targeted companies. In two separate incidents, the most recent of which occurred in 2025, individuals used fraudulent identification in connection with their hiring by the Company. While these individuals were detected and terminated before any data was extracted or malware installed, there can be no assurance that future attempts by similar actors will be unsuccessful.

System vulnerabilities and/or cybersecurity breaches experienced by our third-party service providers have constituted a substantial share of the information security risks that have affected us. For example, in the past we have experienced cybersecurity incidents involving third-party service providers, including incidents in which unauthorized third parties accessed or exfiltrated certain information, including non-significant Amgen data and personally identifiable patient information, and we

have made required regulatory notifications in connection with certain such incidents. In November 2025, a third-party service provider had a cybersecurity incident in which Amgen-related information was accessed and disclosed by a threat actor that, upon review, did not involve information material to the Company. Although these supplier data breaches have not resulted in material adverse effects on our business, there can be no assurance that a similar future cybersecurity incident would not result in a material adverse effect on our business or results of operations. Further, the timeliness of our awareness of a cybersecurity incident affects our ability to respond to and work to mitigate the severity of such events. For example, we have previously experienced incidents in which third-party vendors initially reported that cyberattacks did not involve our data but later determined that attackers had accessed limited, non-significant Amgen information. Although such incidents have not resulted in significant adverse effects on our business, future incidents in which we do not receive timely or complete information regarding the nature or scope of a cybersecurity event could impair our response and could have a material adverse effect on our business.

Cyberattackers are also increasingly exploiting vulnerabilities in commercially available software from shared or open-source code. We rely on third party commercial software that have had and may have such vulnerabilities, but as use of open-source code is frequently not disclosed, our ability to fully assess this risk to our systems is limited. Although these vulnerabilities did not result in any significant adverse effects on us, there can be no assurances that a similar future vulnerability in the software and services that we use would not result in a material adverse effect on our business or results of operations.

Domestic and global government regulators, our business partners, suppliers with whom we do business, companies that provide us or our partners with business services and companies we have acquired or may acquire face similar risks. Security breaches of their systems or service outages have adversely affected systems and could, in the future, affect our systems and security, leave us without access to important systems, products, raw materials, components, services or information, or expose our confidential data or sensitive personal information. For example, we have experienced cybersecurity incidents at third-party vendors that provide testing, analytical, information technology and clinical data services, which required us to temporarily disconnect our systems from those vendors. Although these incidents did not result in breaches of our systems or significantly affect product availability, a prolonged or more widespread service outage affecting these or other vendors, particularly where a vendor is a single source for critical services, could have a material adverse effect on our business or results of operations. In 2024, Change Healthcare, a large U.S. insurance claim and co-pay card processing clearinghouse, experienced a ransomware attack that has caused significant disruptions to healthcare provider and pharmacy operations. While Change Healthcare does not directly provide us with services, disruptions to co-pay card support, insurance billing and Medicaid rebate processing led to lost sales and required us to take action to help patients access their medications and to provide extended payment terms to certain customers. Although services have been rerouted and restored, and the impact on our business was immaterial, similar disruptions may occur in the future stemming from the interconnectedness of the U.S. healthcare ecosystem and industry reliance on centralized claims processing systems and networks, and such future disruptions may have a material adverse effect on our business or results of operations. In addition, we distribute our products in the United States primarily through three pharmaceutical wholesalers, and a security breach that impairs the distribution operations of our wholesalers could significantly impair our ability to deliver our products to healthcare providers and patients. There can be no assurance that our cybersecurity risk management program and processes, including our policies, controls, or procedures, will be fully implemented, complied with or effective in protecting our information technology systems and sensitive data.

Although we have experienced system breakdowns, attacks and information security breaches, we do not believe such breakdowns, attacks and breaches have had a material adverse effect on our business or results of operations. We will continue to experience varying degrees of cyberattacks and other incidents in the future. Even though we continue to invest in the monitoring, protection and resilience of our critical and/or sensitive data and systems, there can be no assurance that our efforts will detect, prevent or fully recover systems or data from all breakdowns, service interruptions, attacks and/or breaches of our systems that could adversely affect our business and operations and/or result in the loss or exposure of critical, proprietary, private, confidential or otherwise sensitive data, which could result in material financial, legal business or reputational harm to us or negatively affect our stock price. While we maintain cyber-liability insurance, our insurance is not sufficient to cover us against all losses that could potentially result from a service interruption, breach of our systems or loss of our critical or sensitive data.

We are also subject to various laws and regulations globally regarding cybersecurity, privacy and data protection, including laws and regulations relating to the collection, storage, handling, use, disclosure, transfer and security of personal data. The legislative and regulatory environment regarding privacy and data protection continues to evolve and receive significant attention globally. For example, we are subject to the EU's GDPR and the California Consumer Privacy Act, as amended (CCPA), both of which impose comprehensive data protection obligations and provide for substantial penalties for noncompliance. Similar consumer privacy and data protection laws have been enacted or proposed in more than half of U.S. states, many of which impose obligations and restrictions that are comparable to, or in some cases more stringent than, those under the CCPA. Outside the United States, other jurisdictions where we operate have passed, or continue to propose, data

privacy or cybersecurity legislation and/or regulations. For example, in China, the Personal Information Protection Law and the Data Security Law, which regulate data processing activities associated with personal and nonpersonal data, builds upon the existing Cybersecurity Law. Failure to comply with these current and future laws could result in significant penalties and reputational harm and could have a material adverse effect on our business and results of operations.

We are adopting and exploring the use of AI in our business, and as an emerging and rapidly evolving technology, our use of AI introduces potential opportunities but also presents risks that could adversely affect our operations, information security and reputation. AI systems may produce inaccurate or flawed outputs due to flawed algorithms, or insufficient and/or erroneous training data. Reliance on flawed outputs could prevent us from effectively utilizing AI in our business or result in lower quality decision-making. We may also become vulnerable to operational disruptions if the AI technologies we use experience downtimes or are compromised by cyberattacks. If we do not effectively implement guardrails and train our staff on the safe and proper use of AI, or if our staff fail to effectively adhere to our established guardrails and training on the use of AI, we may experience adverse effects on our business, including data breaches, the loss of confidential information (including our intellectual property), unintentional disclosure of personal data, or other misuse of our proprietary information. Further, several governments and regulatory authorities have proposed or passed laws and regulations governing the use of AI. For example, the European Parliament has adopted the Artificial Intelligence Act establishing EU-wide rules on data quality, transparency, human oversight and accountability with respect to the use of AI, and U.S. federal and state governments, including California, have enacted or are considering laws regulating the development and use of AI. In 2024, the EU also revised its Cybersecurity Directive NIS2 rules that create new cybersecurity risk management and reporting obligations. Failure to comply with these current and future laws could result in significant penalties and reputational harm and could have a material adverse effect on our business and results of operations.

Our sales and operations are subject to the risks of doing business internationally, including in new or emerging markets.

As we continue our expansion efforts in emerging markets around the world, through acquisitions and licensing transactions as well as through the development and introduction, both independently and through collaborations such as our collaboration with BeOne, of our products in new markets, we face numerous risks to our business. There is no guarantee that our efforts and strategies to expand sales in new or emerging markets will succeed. Our international business, including in China and emerging market countries, may be especially vulnerable to periods of global, national, and local political, legal, regulatory and financial instability, including issues of geopolitical relations, the imposition of international sanctions in response to certain state actions and/or sovereign debt issues, and management of health and healthcare policies. For example, the BIOSECURE Act, signed into law in December 2025 and subject to phased implementation over the next several years through agency guidance and rulemaking, prohibits federal contracting with companies that have commercial connections with enumerated “biotechnology companies of concern” located in certain geographies, including China, could restrict our ability to contract or collaborate with such biotechnology companies in the future. If relations between the United States and other governments deteriorate, our business and investments in such markets may also be adversely affected. We may also be required to increase our reliance on third-party agents and unfamiliar operations and arrangements, including those previously utilized by companies we partner with or acquire in emerging markets. See *We must conduct clinical trials in humans before we commercialize and sell any of our product candidates or existing products for new indications*. Our expansion efforts in China and new and emerging markets around the world are dependent upon the existence or establishment of an environment that is predictable, navigable and supportive of biopharmaceutical innovation, sustained access for our products and predictable pricing controls. China has implemented and enforced regulations governing human genetic resources, including strict requirements with respect to the collection, use and transmission of Chinese human genetic materials and data, and has expanded regulations on the conduct of biotechnology R&D activities in China. For example, between 2020 and 2022, we experienced delays in our applications to the Human Genetic Resources Administration of China that sought approval to conduct clinical trials in China. Further, recent increases in tariffs imposed on certain goods imported into the United States, including inputs relevant to biopharmaceutical manufacturing, have raised our production costs to a limited degree in 2025, and, going forward, such tariffs, together with the imposition of tariffs from agreements the Administration has made with other countries and other potential future tariffs, could further increase our production costs and/or potentially disrupt the operation of our supply chain. See *Global economic conditions may negatively affect us and may magnify certain risks that affect our business*. Our international operations and business may also be subject to less protective intellectual property or other applicable laws, diverse data privacy and protection requirements, changing tax laws and tariffs, trade restrictions or other barriers designed to protect industry in the home country against foreign competition, far-reaching antibribery and anticorruption laws and regulations and/or evolving legal and regulatory environments. For example, cross-border data transfer compliance requirements in China, as well as the DOJ final rule on preventing access to Americans’ bulk sensitive personal data by “countries of concern,” may also impose additional costs of doing business, including costs associated with localizing operations.

In response to the ongoing armed conflict in Ukraine, the U.S. government, numerous state governments, the EU and other countries in which we conduct business have imposed a wide range of economic sanctions that restrict commerce and

business dealings with Russia, certain regions of Ukraine and certain entities and individuals. Additionally, the recent armed conflict in the Middle East has caused regional disruptions to economic activity. For a description of the conflict's impact on our third-party contract manufacturing of KRYSTEXXA, see *Our efforts to collaborate with or acquire other companies, products, or technology, and to integrate the operations of companies or to support the products or technology we have acquired, may not be successful, and may result in unanticipated costs, delays or failures to realize the benefits of the transactions.* These conflicts, in addition to other geopolitical tensions, may also precipitate or amplify the other risks described herein, including risks relating to global economic conditions, cybersecurity, clinical trials and supply chains, which could adversely affect our business, operations and financial condition and results.

We are subject to fluctuations in foreign currency exchange rates relative to the U.S. dollar in the non-U.S. jurisdictions where we do business. While we have a program in place that is designed to reduce our exposure to foreign currency exchange rate fluctuations through foreign currency hedging arrangements, our hedging efforts do not completely offset the effect of these fluctuations on our revenues and earnings. Overall, the legal and operational challenges of our international business operations, along with government controls, the challenges of attracting and retaining qualified personnel and obtaining and/or maintaining necessary regulatory or pricing approvals of our products, may result in material adverse effects on our international product sales, business and results of operations. In addition, pricing pressures and price controls in non-U.S. jurisdictions could adversely affect our sales and revenues, and further, to the extent that those prices are included in MFN calculations, such inclusion may affect our strategy with respect to such jurisdictions.

We may not be able to access the capital and credit markets on terms that are favorable to us, or at all.

The capital and credit markets may experience extreme volatility and disruption, which may lead to uncertainty and liquidity issues for both borrowers and investors. For example, in early 2020, there were significant disruptions in the commercial paper market and several borrowers were unable to obtain funding at normal rates or maturities, which resulted in a significant increase in draws of corporate credit lines with banks. Similarly, the bond markets experienced extreme volatility in terms of interest rates and credit spreads, with several days without new issuances of corporate bonds.

We expect to access the capital markets, from time to time, to supplement our existing funds and cash generated from operations to satisfy our needs for working capital; capital expenditure and debt service requirements; our plans to pay dividends and repurchase stock; and other strategic business initiatives we plan to pursue, including acquisitions and licensing activities. In the event of adverse capital and credit market conditions, we may be unable to obtain capital market financing on favorable terms, or at all, which could have a material adverse effect on our business and results of operations or our ability to complete business acquisitions. Changes in credit ratings issued by nationally recognized credit-rating agencies could also adversely affect our ability to obtain capital and credit market financing and the cost of such financing and have an adverse effect on the market price of our securities.

RISKS RELATED TO COMPETITION

Our products face substantial competition and our product candidates are also likely to face substantial competition.

We operate in a highly competitive environment. See Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Competition*. We expect that our products and product candidates will compete with existing drugs, new drugs currently in development, drugs currently approved for other indications that may later be approved for the same indications as those of our products and drugs approved for other indications that are used off-label. Large pharmaceutical companies and generics manufacturers of pharmaceutical products have expanded into, and are expected to continue expanding into, the biotechnology field, and some pharmaceutical companies and generics manufacturers have formed partnerships to pursue biosimilars. With the proliferation of companies pursuing biopharmaceuticals, several of our biosimilar products have entered, and a number of our product candidates are expected to enter, markets with one or more competitors or with competitors soon to arrive. In addition, some of our competitors may have technical, competitive or other advantages over us for the development of technologies and processes or greater experience in particular therapeutic areas, and consolidation among pharmaceutical and biotechnology companies can enhance such advantages. These advantages may make it difficult for us to compete with them successfully to discover, develop and market new products and for our current products to compete with new products or new product indications they may bring to market. As a result, our products have been competing and may continue to compete, and our product candidates may compete, against products or product candidates that offer higher rebates or discounts, lower prices, equivalent or superior efficacy, better safety profiles, easier administration, earlier market availability, established market position or other competitive features. If we are unable to compete effectively, this could reduce our sales, which could have a material adverse effect on our business and results of operations.

Our intellectual property positions may be challenged, invalidated or circumvented, or we may fail to prevail in current and future intellectual property litigation.

Our success depends in part on our ability to obtain and defend patent rights and other intellectual property rights that are important to the commercialization of our products and product candidates. The patent positions of pharmaceutical and biotechnology companies can be highly uncertain and often involve complex legal, scientific and factual questions. Driven by cost pressures, efforts to limit or weaken patent protection for our industry are increasing. For example, the COVID-19 pandemic resulted in increased interest in compulsory licenses, march-in rights or other governmental interventions, both in the United States and internationally, related to the procurement of drugs, and governments and international bodies have since adopted or considered regulations that could facilitate the use of such measures during public health or other emergencies. At the end of 2023, the prior Administration released a proposed framework that would consider price as a factor when determining whether to exercise march-in rights pursuant to the Bayh-Dole Act with respect to drugs or other taxpayer-funded inventions, and the status and potential implementation of such framework remains uncertain. Further, in early 2025, the Administration took actions affecting federal workforce staffing and operations. Significant reductions of, or disruptions to, staffing and resources available at the USPTO could lead to delays in the examination or approval of patent applications, or other challenges to securing and/or enforcing our intellectual property rights.

Third parties have challenged and may continue to challenge, invalidate or circumvent our patents (including any patent applications, term extensions, term adjustments and supplemental protection certificates) relating to our products, product candidates and technologies. See Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements. Challenges to patents have come from potential competitors or from parties other than those who sought to market a potentially-infringing product. In addition, in certain cases our patent positions have not protected us, and may not protect us, against competitors with similar products or technologies because competing products or technologies may not infringe our patents. For certain of our products and/or product candidates, third parties may obtain patents that claim the product, one or more of its uses, the drug delivery device used to administer the product, or our manufacturing process used to make the product, and they may seek to prevent us from commercializing the product and/or seek payment of a royalty on the product's sales in one or more territories. Further, disputes may arise with third parties from whom we have licensed rights to intellectual property necessary for the development and commercialization of some of our products. In addition, our intellectual property licensing and related business practices have been, and may in the future be, subject to antitrust and competition law scrutiny, which could result in litigation, enforcement actions and/or restrictions on our business. Patent disputes are frequent, costly and can preclude, delay or increase the cost of commercialization of products. We have been in the past, are currently and expect to be in the future, involved in patent litigation. These matters have included, and may in the future include, litigation with manufacturers of products that purport to be biosimilars of certain of our products for patent infringement, invalidity, unenforceability and failure to comply with certain provisions of the BPCIA, and litigation with manufacturers of innovator products that allege patent infringement. A determination made by a court, agency or tribunal concerning infringement, validity, enforceability, injunctive or economic remedy, or the right to patent protection, for example, are typically subject to appellate or administrative review. Upon review, such initial determinations may be afforded little or no deference by the reviewing tribunal and may be affirmed, reversed or made the subject of reconsideration through further proceedings. A patent dispute or litigation has not discouraged, and may not in the future discourage, a potential violator from bringing the allegedly infringing product to market prior to a final resolution of the dispute or litigation. The period from inception until resolution of a patent dispute or litigation is subject to the availability and schedule of the court, agency or tribunal before which the dispute or litigation is pending. We have been, and may in the future be, subject to competition during this period and may not be able to recover fully from the losses, damages and harms we incur from infringement by the competitor product even if we prevail. Moreover, if we lose or settle current or future litigations at certain stages or entirely, we could be subject to competition and/or significant liabilities, be required to enter into third-party licenses for the infringed product or technology or be required to cease using the technology or product in dispute. In addition, we cannot guarantee that such licenses will be available on terms acceptable to us, or at all.

Further, under the Hatch-Waxman Act, our products approved by the FDA under the FDCA have been, and may in the future be, the subject of patent litigation with generics competitors before expiry of the five-year period of data exclusivity provided for under the Hatch-Waxman Act and prior to the expiration of the patents listed for the product. Likewise, our innovative biologic products have been, and may in the future be, the subject of patent litigation prior to the expiration of our patents and, with respect to competitors seeking approval as a biosimilar or interchangeable version of our products, prior to the 12-year exclusivity period provided under the BPCIA. In addition, we have faced, and may in the future face, patent litigation involving claims that our biosimilar product candidates infringe the patents of other companies, including those that manufacture, market or sell the applicable reference products or who are developing or have developed other biosimilar versions of such products. Patents held by other entities have contributed, and may in the future contribute, to a decision by us to not pursue all of the labeled indications of the applicable reference product. However, a decision not to pursue all of the labeled indications of the applicable reference product might not avoid, or end, potential litigation. While we have attempted, and expect to continue to attempt, to challenge the patents held by other companies, our efforts may be unsuccessful. For

examples of and information related to our patent litigation, see Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements.

Certain of the existing patents on our products have expired or will soon expire. See Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Patents*. As our patents expire, competitors are able to legally produce and market similar products or technologies, including biosimilars, which has had, and may continue to have, a material adverse effect on our product sales, business and results of operations. In addition, competitors have been, and may continue to be, able to invalidate, design around or otherwise circumvent our patents and sell competing products.

We currently face competition from biosimilars and generics and expect to face increasing competition from biosimilars and generics in the future.

We currently face competition from biosimilars and generics in most of the territories in which we operate, including the United States and Europe, and we expect to face increasing biosimilar and/or generics competition this year and beyond. Expiration or successful challenge of applicable patent rights or expiration of an applicable exclusivity period has accelerated such competition, and we expect to face more litigation regarding the validity and/or scope of our patents. Our products have also experienced greater competition from lower cost biosimilars or generics that come to market when branded products that compete with our products lose their own patent protection. To the extent that governments adopt more permissive regulatory approval standards and competitors are able to obtain broader or expedited marketing approval for biosimilars and generics, the rate of increased competition for our products would likely accelerate. For example, both the EMA and FDA have increasingly supported streamlined biosimilar approval pathways in which comparative clinical efficacy testing may not be routinely required, and in recent years have, on a case by case basis, accepted applications without such testing.

In the EU, biosimilars are evaluated for marketing authorization pursuant to a set of general and product class-specific guidelines. In addition, in an effort to spur biosimilar utilization and/or increase potential healthcare savings, some EU countries and some Canadian provinces have adopted biosimilar uptake measures such as physician prescribing quotas or automatic pharmacy substitution of biosimilars for the corresponding reference products. Some EU countries impose automatic price reductions upon market entry of one or more biosimilar competitors. Regulatory authorities in the EU, including the EMA and the EU Heads of Medicines' Agencies (HMA) have articulated a position that biosimilar medicines approved in the EU are “interchangeable” with their reference products and other biosimilars of the same reference product for purposes of prescribing. Increased regulatory clarity regarding interchangeability, together with national pricing, tendering and reimbursement policies, has coincided with continued growth in biosimilar utilization and increased competitive pressure on innovative biologic products in Europe. While the degree of competitive effects of biosimilar competition differs between EU countries and between products, biosimilar competition has had, and is expected to continue to have, an effect on the sales of innovative biologic products in Europe.

In the United States, the BPCIA authorizes the FDA to approve biosimilars via a separate, abbreviated pathway. See Item 1. Business—Government Regulation—*Regulation in the United States—Approval of Biosimilars*. In the United States, the FDA has approved numerous biosimilars, including biosimilar versions of Neulasta, EPOGEN, ENBREL, Prolia and XGEVA, and a growing number of companies have announced that they are also developing biosimilar versions of our products. For example, a number of biosimilar versions of Prolia and/or XGEVA have been approved in the United States and EU, and impact to our Prolia and XGEVA sales are expected to accelerate as the number of additional competitors continues to increase. See Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Competition*. Manufacturers of biosimilars have attempted, and may in the future attempt, to compete with our products by offering greater discounts or rebates, contracts that offer longer-term pricing or a broader portfolio of other products, or lower list prices. Companies pursuing development of biosimilar versions of our products have challenged and may continue to challenge our patents well in advance of the expiration of our material patents. For examples of and information related to our biosimilars and generics patent litigation, see Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements. See also *Our intellectual property positions may be challenged, invalidated or circumvented, or we may fail to prevail in current and future intellectual property litigation*.

Under the U.S. biosimilar regulatory framework, certain biosimilars have been approved as interchangeable with their reference products, which can allow pharmacists, subject to state law, to substitute those biosimilars for our products without prior prescriber approval and could increase competitive pressure on our products. In addition, critics of the 12-year exclusivity period for biologics may continue to seek to shorten that period or encourage narrower interpretations of the statutory provisions governing eligibility for exclusivity, and the FDA could adopt interpretations of the biologics exclusivity framework that limit the scope or duration of exclusivity for certain products or product changes. Legislative and regulatory initiatives may seek to further promote competition by facilitating generic and biosimilar approval and commercialization, including by reducing regulatory requirements, limiting the scope of patent protections or disputes, or adopting reimbursement, substitution or disclosure policies that favor biosimilars. Any such developments could accelerate biosimilar competition and adversely affect the sales of our products.

Upon the expiration or loss of patent protection and/or applicable exclusivity for one of our products, we can lose the majority of revenues for that product in a very short period of time. See Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Competition*. Additionally, if one of our products is the subject of an FDA Written Request for pediatric studies and we are unable to adequately complete these studies, we may not obtain the pediatric exclusivity award that extends unexpired regulatory exclusivity for the product (and existing patents for a small molecule product) by an additional six months. Further, the FDA released draft guidance contemplating that pediatric exclusivity may not be granted for studies conducted solely to fulfill Pediatric Research Equity Act (PREA) requirements.

While we are unable to predict the precise effects and timing of biosimilars and generics on our products, we are currently facing and expect to face greater competition in the United States, Europe and elsewhere as a result of biosimilar and generic competition and, in turn, downward pressure on our product prices and sales. This competition has had, and could increasingly have, a material adverse effect on our product sales, business and results of operations. State laws may also have an impact on our business. For example, certain states have enacted, or proposed, laws that restrict, or prohibit, “pay for delay” settlements of patent infringement claims filed by manufacturers of generics or biosimilars by creating the presumption that such agreements are anticompetitive. In addition, federal legislative proposals targeting such settlements continue to be introduced and advanced in Congress. These developments could result in prolonged litigation or fewer settlements of patent infringement disputes.

Concentration of sales at certain of our wholesaler distributors, and consolidation of private payers, such as insurers, and PBMs has negatively affected, and may continue to negatively affect, our business.

Certain of our distributors, customers and payers have substantial purchasing leverage, due to the volume of our products they purchase or the number of patient lives for which they provide coverage. The substantial majority of our U.S. product sales is made to three pharmaceutical product wholesaler distributors: McKesson Corporation, Cencora, Inc. (formerly AmerisourceBergen Corporation) and Cardinal Health, Inc. These distributors, in turn, sell our products to their customers, which include physicians or their clinics, dialysis centers, hospitals and pharmacies. Similarly, as discussed above, there has been significant consolidation in the health insurance industry, including that a small number of PBMs now oversee a substantial percentage of total covered lives in the United States. See *Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability*. For example, the six largest PBMs in the United States are now part of major health insurance providers, and the Federal Trade Commission’s interim report released in 2024 showed that they nationally account for 94% of prescription drug claims. The growing concentration of purchasing and negotiating power by these entities has, and may continue to, put pressure on our pricing due to their ability to extract price discounts on our products, fees for other services or rebates, negatively affecting our bargaining position, sales and/or profit margins. In addition, decisions by these entities to purchase or cover less or none of our products in favor of competing products could have a material adverse effect on our product sales, business and results of operations due to their purchasing volume. Further, if one of our significant wholesale distributors encounters financial or other difficulties and becomes unable or unwilling to pay us all amounts that such distributor owes us on a timely basis, or at all, it could negatively affect our business and results of operations. In addition, if one of our significant wholesale distributors becomes insolvent or otherwise unable to continue its commercial relationship with us in its present form, it could significantly disrupt our business and adversely affect our product sales, our business and results of operations unless suitable alternatives are timely found or lost sales are absorbed by another distributor.

RISKS RELATED TO RESEARCH AND DEVELOPMENT

We may not be able to develop commercial products despite significant investments in R&D.

Amgen invests heavily in R&D. Successful product development in the biotechnology industry is highly uncertain, and very few R&D projects yield approved and commercially viable products. Product candidates, including biosimilar product candidates, or new indications for existing products (collectively, product candidates) that appear promising in the early phases of development have failed to reach the market for a number of reasons, such as:

- the product candidate did not demonstrate acceptable clinical trial results even though it achieved its primary endpoints and/or demonstrated positive preclinical or early clinical trial results, for reasons that could include changes in the standard of care of medicine or expectations of health authorities;
- the product candidate was not effective or not more effective than currently available or potentially competitive therapies in treating a specified condition or illness;
- the product candidate was not cost effective in light of existing or potentially competitive therapeutics;
- the product candidate had harmful side effects in animals or humans;
- the necessary regulatory bodies, such as the FDA or EMA, did not approve the product candidate for an intended use;

- reimbursement for the product candidate is limited despite regulatory approval;
- the product candidate was not economical for us to manufacture and/or commercialize;
- the patient population size is smaller than anticipated;
- other parties had or may have had proprietary rights relating to our product candidate, such as patent rights, and did not let us sell it on reasonable terms, or at all;
- we and certain of our licensees, partners, contracted organizations or independent investigators failed to effectively conduct clinical development or clinical manufacturing activities;
- the pathway to regulatory approval or reimbursement for product candidates was uncertain or not well-defined;
- the biosimilar product candidate failed to demonstrate the requisite biosimilarity to the applicable reference product, or was otherwise determined by a regulatory authority to not meet applicable standards for approval; and
- a companion diagnostic device that is required with the use of a product candidate is not approved by the necessary regulatory authority.

We believe that genetics, together with the benefit of artificial intelligence and computational evidence, could meaningfully aid our search for new medicines and help guide our R&D decisions and investments, and have focused our R&D strategy on drug targets validated by genetic or other compelling human evidence. We have invested considerable time, energy and resources into developing our expertise in human genetics and acquiring access to libraries of genetic information, and are applying artificial intelligence to our R&D activities, including applying such technologies to advance our human data efforts and our generative biology platform that seek to discover and design new drugs. However, product candidates based on genetically validated targets or developed with the assistance of such technologies remain subject to the uncertainties of the drug development process and may not reach the market for a number of reasons, including the factors listed above.

We must conduct clinical trials in humans before we commercialize and sell any of our product candidates or existing products for new indications.

Before a product may be sold, we must conduct clinical trials to demonstrate that our product candidates are safe and effective for use in humans. The results of those clinical trials are used as the basis to obtain approval from regulatory authorities such as the FDA and EMA. See *Our current products and products in development cannot be sold without regulatory approval*. We are required to conduct clinical trials using an appropriate number of trial sites and patients to support the product label claims. The length of time, number of trial sites and number of patients required for clinical trials vary substantially, and we may spend several years and incur substantial expense in completing certain clinical trials. In addition, we may have difficulty finding a sufficient number of clinical trial sites and/or patients to participate in our clinical trials, particularly if competitors are conducting clinical trials in similar patient populations and/or in rare disease therapy clinical trials due to the inherently small patient population potentially served by such therapies. Patients may withdraw from clinical trials at any time (including trials in which patients believe that they may not be receiving a clinical benefit), and evolving legal obligations including, but not limited to, privacy laws and/or other restrictions in certain countries may restrict the ability of clinical trial investigators to conduct further follow-up on such patients, which may adversely affect the interpretation of study results. Regulatory authorities may also pause or halt conduct of clinical trials based on their appraisal of the potential or actual risks of continuing the study. Delays and complications in planned clinical trials can result in increased development costs, associated delays in regulatory approvals and in product candidates reaching the market and revisions to existing product labels.

Further, to increase the number of patients available for enrollment in our clinical trials, we have opened, and will continue to open, clinical sites and enroll patients in a number of locations where our experience conducting clinical trials is more limited, including India, China, South Korea, the Philippines, Singapore, Saudi Arabia and some Central and South American countries, either through utilization of third-party contract clinical trial providers entirely or in combination with local staff. Conducting clinical trials in locations where we have limited experience requires substantial time and resources to understand the unique regulatory environments of individual countries. For other examples of the risks of conducting clinical trials in China, see also *Our sales and operations are subject to the risks of doing business internationally, including in emerging markets*. Further, we must ensure the timely production, distribution and delivery of the clinical supply of our product candidates to numerous and varied clinical trial sites. Additionally, regional disruptions, including natural and man-made disasters, health emergencies (such as novel viruses or pandemics), or geopolitical conflicts (such as the ongoing armed conflicts in Ukraine and the Middle East) have significantly disrupted the timing of clinical trials, and in the future could disrupt the timing, execution and outcome of clinical trials. If we fail to adequately manage the design, execution and diverse regulatory aspects of our clinical trials or to manage the production or distribution of our clinical supply, or such sites experience disruptions as a result of a natural/man-made disaster, health emergency or geopolitical conflict, corresponding

regulatory approvals may be delayed or we may fail to gain approval for our product candidates or could lose our ability to market existing products in certain therapeutic areas or altogether. If we are unable to market and sell our products or product candidates or to obtain approvals in the timeframe needed to execute our product strategies, our business and results of operations could be materially and adversely affected.

We rely on independent third-party clinical investigators to recruit patients and conduct clinical trials on our behalf in accordance with applicable study protocols, laws and regulations. We also rely on unaffiliated third-party vendors to perform certain aspects of our clinical trial operations, including that such vendors have appropriate experienced staff to execute on such activities. Further, the disease states that we are studying, such as cancers, require complex treatment protocols that may be difficult to consistently apply across global trial sites, which can impact the quality, interpretability, timing and/or registrability of the data generated. In some circumstances, we enter into co-development arrangements with other pharmaceutical and medical devices companies that provide for the other company to conduct certain clinical trials for the product we are co-developing or to develop a diagnostic test used in screening or monitoring patients in our clinical trials. See *Some of our pharmaceutical pipeline and our commercial product sales rely on collaborations with third parties, which may adversely affect the development and sales of our products*. We also may acquire companies that have past or ongoing clinical trials or rights to products or product candidates for which clinical trials have been or are being conducted. These trials may not have been conducted to the same standards as ours; however, once an acquisition has been completed we assume responsibility for the conduct of these trials, including any potential risks and liabilities associated with the past and prospective conduct of those trials. If regulatory authorities determine that we or others, including our licensees or co-development partners, or the independent investigators or vendors selected by us, our co-development partners or by a company we have acquired or from which we have acquired rights to a product or product candidate, have not complied with regulations applicable to the clinical trials, those authorities may refuse or reject some or all of the clinical trial data or take other actions that could delay or otherwise negatively affect our ability to obtain or maintain marketing approval of the product or indication. In addition, delays or failures to develop diagnostic tests or delivery devices for our clinical trials can affect the timely enrollment of such trials and lead to delays or inability to obtain marketing approval. If we were unable to market and sell our products or product candidates, our business and results of operations could be materially and adversely affected.

In addition, some of our clinical trials utilize drugs and combination products manufactured and marketed by other pharmaceutical companies or vendors. These drugs, devices and/or products may be administered or used in clinical trials in combination with one of our products or product candidates or in a head-to-head study comparing the products' or product candidates' relative efficacy and safety. In the event that any of these vendors or pharmaceutical companies have unforeseen issues that negatively affect the quality of their work product or create a shortage of supply, or if we are otherwise unable to obtain an adequate supply of these other drugs, our ability to complete our applicable clinical trials and/or evaluate clinical results may also be negatively affected. As a result, such quality or supply problems could adversely affect our ability to timely file for, gain or maintain regulatory approvals worldwide.

Clinical trials must generally be designed based on the current standard of medical care. However, in certain diseases, such as cancer, the standard of care is evolving rapidly. In some cases, we may design a clinical trial based on the standard of care we anticipate will exist at the time our study is completed. The duration of time needed to complete certain clinical trials may result in the design of such clinical trials being based on standards of medical care that are no longer or that have not become the current standards by the time such trials are completed, limiting the utility and application of such trials. Additionally, the views of regulatory agencies relating to the requirements for accelerated approval have evolved over time, and trial designs that were sufficient to support accelerated approvals for some oncology products may not be considered sufficient for later candidates. In addition, access to expedited regulatory programs, including priority review vouchers and other programs tied to agency-designated national or public health priorities, is subject to evolving agency eligibility criteria and guidance, and may not be available for our product candidates or may not result in expedited review or approval. We may not obtain favorable clinical trial results and therefore may not be able to obtain regulatory approval for new product candidates or new indications for existing products and/or maintain our current product labels. Participants in clinical trials of our products and product candidates may also suffer adverse medical events or side effects that could, among other factors, delay or terminate clinical trial programs and/or require additional or longer trials to gain approval.

Even after a product is on the market, safety concerns may require additional or more extensive clinical trials as part of a risk management plan for our product or for approval of a new indication. Additional clinical trials we initiate, including those required by the FDA, could result in substantial additional expense, and the outcomes could result in further label restrictions or the loss of regulatory approval for an approved indication, each of which could have a material adverse effect on our product sales, business and results of operations. Additionally, any negative results from such trials could materially affect the extent of approvals, the use, reimbursement and sales of our products, our business and results of operations.

Our current products and products in development cannot be sold without regulatory approval.

Our business is subject to extensive regulation by numerous state and federal government authorities in the United States, including the FDA, and by foreign regulatory authorities, including the EMA. We are required in the United States and in the other regions and countries in which we, or our partners and affiliates, sell to obtain approval from regulatory authorities before we manufacture, market and sell our products. Once our products are approved, the FDA and other U.S. and ex-U.S. regulatory agencies have substantial authority to require additional testing and reporting, perform inspections, change product labeling or mandate withdrawals of our products. Failure to comply with applicable regulatory requirements may subject us to administrative and/or judicially imposed sanctions or monetary penalties as well as reputational and other harms. The sanctions could include the FDA's or ex-U.S. regulatory authorities' refusals to approve pending applications, delays in obtaining or withdrawals of approvals, delays or suspensions of clinical trials, warning letters, product recalls or seizures, total or partial suspensions of our operations, injunctions, fines, civil penalties and/or criminal prosecutions.

Obtaining and maintaining regulatory approvals have been, and will continue to be, increasingly difficult, time-consuming and costly. Legislative bodies or regulatory agencies could enact new laws or regulations, change existing laws or regulations or change their interpretations of laws or regulations at any time, which could affect our ability to obtain or maintain approval of our products or product candidates. The rate and degree of change in existing laws and regulations and regulatory expectations have accelerated in established markets, and regulatory expectations continue to evolve in emerging markets. We are unable to predict whether and when any further changes to laws or regulatory policies affecting our business could occur, such as changes to laws or regulations governing manufacturer communications concerning drug products and drug product candidates and whether such changes could have a material adverse effect on our product sales, business and results of operations. Further, we are reliant on regulators having the resources necessary to evaluate and approve our products. In the United States, federal government shutdowns have occurred and such shutdowns have disrupted certain regulatory activities. Any future government shutdowns or workforce constraints may delay or disrupt regulatory activities, including those with respect to our ongoing clinical programs, the manufacture of our products and product candidates and product approvals.

Recent initiatives to reduce the size and budgets of government agencies, including the HHS, FDA and NIH, may adversely impact our operations. In particular, reductions in staffing and resources at the FDA could result in delays in regulatory review timelines and marketing application and supplement approvals. Changes in leadership at the FDA may also result in shifts in perspectives on the drug approval process and regulatory priorities. Further, implementation of new policy initiatives without guidance or rulemaking has reduced transparency and increased uncertainty with respect to agency actions. Additionally, funding reductions and caps on research overhead costs imposed on the NIH and its programs may result in grant funding cutbacks for scientific and disease-related research at academic institutions and research centers, and such reductions, over the longer term, may slow the overall discovery and development of new therapies and/or slow or interrupt the flow of innovation into the pharmaceutical development pipeline. These developments and others associated with the reduction of personnel and budgets at the regulatory agencies that oversee our industry and operations may adversely affect our business activities, including our ongoing and future clinical research and drug development programs, research collaborations, manufacturing activities and regulatory submissions.

Regulatory authorities have questioned, and may in the future question, the sufficiency for approval of the endpoints we select for our clinical trials. A number of our products and product candidates have been evaluated in clinical trials using surrogate endpoints that measure an effect that is known to correlate with an ultimate clinical benefit. For example, a therapeutic oncology product candidate may be evaluated for its ability to reduce or eliminate minimal residual disease (MRD), or to extend the length of time during and after the treatment that a patient lives without the disease worsening, measured by progression-free survival (PFS). Demonstrating that the product candidate induces MRD-negative responses or produces a statistically significant improvement in PFS does not necessarily mean that the product candidate will show a statistically significant improvement in overall survival or the time that the patients remain alive. In the cardiovascular setting, a heart disease therapeutic candidate may be evaluated for its ability to reduce LDL-C levels, as an elevated LDL-C level has been a surrogate endpoint for cardiovascular events such as death, heart attack and stroke. The use of surrogate endpoints such as PFS and LDL-C reduction, in the absence of other measures of clinical benefit, may not be sufficient for broad usage or approval even when such results are statistically significant. Regulatory authorities could also add new requirements, such as the completion of enrollment in a confirmatory study or the completion of an outcomes study or a meaningful portion of an outcomes study, as conditions for obtaining approval or obtaining an indication. For example, despite demonstrating that Repatha reduced LDL-C levels in a broad patient population, only after our large phase 3 outcomes study evaluating the ability of Repatha to prevent cardiovascular events met certain of its primary composite endpoint and key secondary composite endpoint did the FDA grant a broader approval of Repatha to reduce the risk of certain cardiovascular events. There may also be situations in which demonstrating the efficacy and safety of a product candidate may not be sufficient to gain regulatory approval unless superiority to other existing treatment options can be shown. The imposition of additional requirements or our inability to meet them in a timely fashion, or at all, has delayed, and may in the future delay, our clinical development and

regulatory filing efforts, delay or prevent us from obtaining regulatory approval for new product candidates or new indications for existing products, or prevent us from maintaining our current product labels.

Some of our products have been approved by U.S. and ex-U.S. regulatory authorities on an accelerated or conditional basis with full approval conditioned upon fulfilling the requirements of regulators. For example, the FDA has approved LUMAKRAS under accelerated approval for the treatment of adult patients with KRAS G12C-mutated local advanced or metastatic NSCLC. Following our submission of the LUMAKRAS/LUMYKRAS CodeBreaK 200 Phase 3 confirmatory data in March 2023 to the FDA and EMA, we received a Complete Response Letter from the FDA and a new post-marketing requirement for an additional confirmatory study to support full approval. Regulatory authorities are placing greater focus on whether the sponsors of products originally approved on an accelerated or conditional basis have met the conditions of the accelerated or conditional approvals. If we are unable to fulfill the regulators' requirements that were conditions of a product's accelerated or conditional approval and/or if regulators reevaluate the data or risk-benefit profile of our product, the conditional approval may not result in full approval or may be revoked or not renewed. Alternatively, we may be required to change the product's labeled indications, conduct an additional confirmatory clinical trial, or even withdraw the product from the market.

Regulatory authorities can also impose post-marketing pediatric study requirements. Failure to fulfill such requirements may result in regulatory or enforcement action, including financial penalties or the invalidation of a product's marketing authorization.

Safety problems or signals can arise as our products and product candidates are evaluated in clinical trials, including investigator sponsored studies, or as our marketed products are used in clinical practice. We are required continuously to collect and assess adverse events reported to us and to communicate to regulatory agencies these adverse events and safety signals regarding our products. Regulatory agencies periodically perform inspections of our pharmacovigilance processes, including our adverse event reporting. In the United States, for our products with approved Risk Evaluation and Mitigation Strategies (REMS, see Item 1. Business—Government Regulation—*Post-approval Phase*), we are required to submit periodic assessment reports to the FDA to demonstrate that the goals of the REMS are being met. REMS and other risk management programs are designed to help ensure that a drug's benefits outweigh the risks and vary in the elements they contain. If the FDA is not satisfied with the results of the periodic assessment reports we submit for any of our REMS, the FDA may also modify our REMS or take other regulatory actions, such as implementing revised or restrictive labeling. The drug delivery devices approved for use in combination with our products are also subject to regulatory oversight and review for safety and malfunctions. See *Some of our products are used with drug delivery or companion diagnostic devices that have their own regulatory, manufacturing and other risks*. If regulatory agencies determine that we or other parties (including our clinical trial investigators, those operating our patient support programs or licensees of our products) have not complied with the applicable reporting, other pharmacovigilance or other safety or quality assessment requirements, we may become subject to additional inspections, warning letters or other enforcement actions, including fines, marketing authorization withdrawal and other penalties. Our product candidates and marketed products can also be affected by safety problems or signals occurring with respect to products that are similar to ours or that implicate an entire class of products. Further, as a result of clinical trials, including sub-analyses or meta-analyses of earlier clinical trials (a meta-analysis involves the use of various statistical methods to combine results from previous separate but related studies) performed by us or others, concerns may arise about the sufficiency of the data or studies underlying a product's approved label. Such actual or perceived safety problems or concerns can lead to:

- revised or restrictive labeling for our products, or the potential for restrictive labeling that has resulted, and may in the future result, in our decision not to commercialize a product candidate;
- requirement of risk management or minimization activities or other regulatory agency compliance actions related to the promotion and sale of our products;
- post-marketing commitments, mandated post-marketing requirements or pharmacovigilance programs for our approved products;
- product recalls of our approved products;
- required changes to the processes used in the manufacture of our products, which could increase our manufacturing costs and affect the availability of contract manufacturers we may utilize to assist in such manufacturing;
- revocation of approval for our products from the market completely, or within particular therapeutic areas or patient types;
- increased timelines or delays in being approved by the FDA or other regulatory bodies; and/or
- treatments or product candidates not being approved by regulatory bodies.

For example, after an imbalance in positively adjudicated cardiovascular serious adverse events was observed in one of the phase 3 clinical trials for EVENITY but not in another, larger phase 3 study, in April 2019 the FDA approved EVENITY for the treatment of osteoporosis in postmenopausal women at high risk for fracture, along with a post-marketing requirement. The requirement includes a five-year observational feasibility study that could be followed by a comparative safety study or trial.

Regulatory authorities also require that our products are tested and controlled for impurities. Impurities exceeding established limits may lead to delayed product approvals or disrupt the manufacture and distribution of our products. For example, certain jurisdictions and regulatory agencies, including the FDA and EMA, require risk assessments, and if applicable, testing, for the presence of nitrosamine impurities in certain small molecule drugs, and we are following the established process of evaluating potentially impacted small molecule products.

In addition to our innovative products, we are working to develop and commercialize biosimilar versions of a number of products currently manufactured, marketed and sold by other pharmaceutical companies. In some markets outside the United States and EU, there is not yet a legislative or regulatory pathway for the approval of biosimilars. In the United States, the BPCIA provides for such a pathway. Discussions within the FDA and other regulatory authorities, and between regulatory authorities and sponsors, continue as to the evidence needed to demonstrate biosimilarity or interchangeability for specific products. See *We currently face competition from biosimilars and generics and expect to face increasing competition from biosimilars and generics in the future*. Delays or uncertainties in the development or implementation of such pathways, or changes in existing regulatory pathways, including degradation of regulatory standards, could result in delays or difficulties in getting our biosimilar products approved by regulatory authorities, subject us to unanticipated development costs or otherwise reduce the value of the investments we have made in the biosimilars area. Further, we cannot predict the extent to which any potential legislative or policy initiatives would affect the biosimilar pathway or have a material adverse effect on our development of biosimilars, on our marketed biosimilars or on our pursuit of interchangeability designations for any biosimilar. In addition, if we are unable to bring our biosimilar products to market on a timely basis and secure “first-to-market” or other advantageous positions, our future biosimilar sales, business and results of operations could be materially and adversely affected.

Some of our products are used with drug delivery or companion diagnostic devices that have their own regulatory, manufacturing and other risks.

Many of our products and product candidates may be used in combination with a drug delivery device, such as an injector or other delivery system. For example, Neulasta is available as part of the Neulasta Onpro kit, our AutoTouch reusable autoinjector is used with ENBREL Mini single-dose prefilled cartridges, Repatha can be administered with the Repatha SureClick autoinjector, and WEZLANA uses our ConfiPen drug delivery device. In addition, some of our products or product candidates, including many of our oncology product candidates and products, including LUMAKRAS/LUMYKRAS, may also require the use of a companion or other diagnostic device such as a device that determines whether the patient is eligible to use our drug or that helps ensure its safe and effective use. In some regions, including the United States, regulatory authorities may require contemporaneous approval of the companion diagnostic device and the therapeutic product; in others the regulatory authorities may require a separate study of the companion diagnostic device. Our product candidates or expanded indications of our products used with such devices may not be approved or may be substantially delayed in receiving regulatory approval if development or approval of such devices is delayed, such devices do not also gain or maintain regulatory approval or clearance, or if such devices do not remain commercially available. When approval of the product and device is sought under a single marketing drug application, the increased complexity of the review process may delay receipt of regulatory approval. In addition, some of these devices may be provided by single-source unaffiliated third-party companies. We are dependent on the sustained cooperation and effort of those third-party companies to supply and/or market the devices and, in some cases, to conduct the studies required for approval or clearance by the applicable regulatory agencies. We are also dependent on those third-party companies continuing to meet applicable regulatory or other requirements. See *We rely on third-party suppliers for certain of our raw materials, medical devices and components*. Failure to successfully develop, modify, or supply the devices, delays in or failures of the Amgen or third-party studies, or failure by us or the third-party companies to obtain or maintain regulatory approval or clearance of the devices could result in increased development costs; delays in, or failure to obtain or maintain, regulatory approval; and/or associated delays in a product candidate reaching the market or in the addition of new indications for existing products. We are also required to collect and assess user complaints, adverse events and malfunctions regarding our devices, and actual or perceived safety problems or concerns with a device used with our product can lead to regulatory actions and adverse effects on our products. See *Our current products and products in development cannot be sold without regulatory approval*. Additionally, regulatory agencies conduct routine monitoring and inspections to identify and evaluate potential issues with our devices. For example, in 2017, the FDA reported on its adverse event reporting system that it was evaluating our Neulasta Onpro kit. Subsequently, we implemented device and labeling enhancements to address product complaints received on this device. We continuously monitor complaints and adverse events and implement additional enhancements as needed. Loss of regulatory approval or clearance of a device that is used with our product may also result in

the removal of our product from the market. Further, failure to successfully develop, supply, or gain or maintain approval for these devices could adversely affect sales of the related approved products. See also *We rely on third-party suppliers for certain of our raw materials, medical devices and components*.

Some of our pharmaceutical pipeline and our commercial product sales rely on collaborations with third parties, which may adversely affect the development and sales of our products.

We depend on alliances with other companies, including pharmaceutical and biotechnology companies, vendors and service providers, for the development of a portion of the products in our pharmaceutical pipeline and for the commercialization and sales of certain of our commercial products. For example, we have collaborations with third parties under which we share development rights, obligations and costs and/or commercial rights and obligations. See Item 1. Business—Business Relationships.

Failures by these parties to meet their contractual, regulatory, or other obligations to us or any disruption in the relationships between us and these third parties, could have a material adverse effect on our pharmaceutical pipeline and business. In addition, our collaborative relationships for R&D and/or commercialization and sales often extend for many years and have given, and may in the future give, rise to disputes regarding the relative rights, obligations and revenues of us and our collaboration partners, including the ownership or prosecution of intellectual property and associated rights and obligations. This could result in the loss of intellectual property rights or protection, delay the development and sale of potential pharmaceutical products, affect the sale and delivery of our commercialized products and lead to lengthy and expensive litigation, administrative proceedings or arbitration.

RISKS RELATED TO OPERATIONS

We perform a substantial majority of our commercial manufacturing activities at our facility in the U.S. territory of Puerto Rico and a substantial majority of our clinical manufacturing activities at our facility in Thousand Oaks, California; significant disruptions or production failures at these facilities could significantly impair our ability to supply our products or continue our clinical trials.

The global supply of our products and product candidates for commercial sales and for use in our clinical trials is significantly dependent on the uninterrupted and efficient operation of our manufacturing facilities, in particular those in the U.S. territory of Puerto Rico and Thousand Oaks, California. See *Manufacturing difficulties, disruptions or delays could limit supply of our products and limit our product sales*.

We currently perform a substantial majority of our clinical manufacturing that supports our product candidates at our facility in Thousand Oaks, California. A substantial disruption in our ability to operate our Thousand Oaks manufacturing facility could materially and adversely affect our ability to supply our product candidates for use in our clinical trials, leading to delays in development of our product candidates.

In addition, we currently perform a substantial majority of our commercial manufacturing activities at our facility in the U.S. territory of Puerto Rico. Puerto Rico has experienced, and continues to be vulnerable to, natural disasters, including hurricanes and earthquakes, that have disrupted the functioning of critical infrastructure such as electric power, water supply, communications networks and transportation systems. The ongoing instability of the island's electric grid has required, at times, sustained reliance on backup generation, and disruptions affecting the island have also impacted certain third-party suppliers and logistics providers. In addition, public health emergencies, labor disruptions, economic challenges and other events affecting Puerto Rico could further disrupt our manufacturing operations, the availability of labor and utilities, and/or the transportation of supplies and finished products. Although our ability to manufacture and supply our products has not, to date, been significantly affected by such events, these challenges, or a combination of these challenges, or other issues that create a substantial disruption to our ability to operate our Puerto Rico manufacturing facility or get supplies and manufactured products transported to and from that location, could make it more expensive or difficult for us to operate in Puerto Rico, and could materially and adversely affect our ability to supply our products and affect our product sales. See *Manufacturing difficulties, disruptions or delays could limit supply of our products and limit our product sales*.

We rely on third-party suppliers for certain of our raw materials, medical devices and components.

We rely on unaffiliated third-party suppliers for certain raw materials, medical devices and components necessary for the manufacturing of our commercial and clinical products. Certain of those raw materials, medical devices and components are proprietary products of those unaffiliated third-party suppliers and are specifically cited in our drug applications with regulatory agencies so that they must be obtained from that specific sole source or sources and could not be obtained from another supplier unless and until the regulatory agency approved such supplier. For example, we rely on a single source for the SureClick autoinjectors used in the drug delivery of a number of our products, including Repatha and ENBREL. Also, certain of the raw

materials required in the commercial and clinical manufacturing of our products are sourced from other countries and/or derived from biological sources, including mammalian tissues, bovine serum and human serum albumin.

Among the reasons we may be unable to obtain these raw materials, medical devices and components include:

- regulatory requirements or action by regulatory agencies or others;
- adverse financial or other strategic developments at, or affecting, the supplier, including bankruptcy;
- unexpected demand for, or shortage of, raw materials, medical devices or components;
- failure to comply with our quality standards which results in quality and product failures, complaints, product contamination and/or recall;
- a material shortage, contamination, recall and/or restrictions on the use of certain biologically derived substances or other raw materials;
- discovery of previously unknown or undetected imperfections in raw materials, medical devices or components;
- cyberattacks on supplier systems;
- natural or other disasters, including hurricanes, earthquakes, volcanoes or fires;
- labor disputes (such as strikes) or shortages, including from the effects of health emergencies (such as novel viruses or pandemics) or natural disasters;
- tariffs or other trade barriers that increase costs, limit availability, or disrupt the flow of goods; and
- geopolitical conflicts (such as the ongoing conflicts in Ukraine and the Middle East).

For example, in prior years we have experienced shortages in certain components necessary for the formulation, fill and finish of certain of our products in our Puerto Rico facility, and we have also experienced shortages related to single use systems and packaging which has caused disruptions to our manufacturing plans. Further quality issues that result in unexpected additional demand for certain components have resulted in shortages and, in the future, may lead to shortages of required raw materials or components. We may experience similar or other shortages in the future resulting in delayed shipments, supply constraints, clinical trial delays, contract disputes and/or stock-outs of our products. These, or other similar events, could negatively affect our ability to satisfy demand for our products or conduct clinical trials, which could have a material adverse effect on our product sales, business and results of operations.

Manufacturing difficulties, disruptions or delays could limit supply of our products and limit our product sales.

Manufacturing biologic and small molecule human therapeutic products is difficult, complex and highly regulated. We manufacture many of our commercial products and product candidates internally. In addition, we use third-party contract manufacturers to produce, or assist in the production of, a number of our products, and we currently use contract manufacturers to produce, or assist in the production of, a number of our late-stage product candidates and drug delivery devices. The number of third-party contract manufacturers that we use has increased with our acquisition of Horizon, as Horizon required contract manufacturers for all of its products. See Item 1. Business—Manufacturing, Distribution and Raw Materials—*Manufacturing*; and *Our efforts to collaborate with or acquire other companies, products, or technology, and to integrate the operations of companies or to support the products or technology we have acquired, may not be successful, and may result in unanticipated costs, delays or failures to realize the benefits of the transactions.* Our ability to adequately and timely manufacture and supply our products (and product candidates to support our clinical trials) is dependent on the uninterrupted and efficient operation of our facilities and those of our third-party contract manufacturers. For example, our acquisition of Horizon resulted in the addition of more than 30 contract manufacturing organizations (CMOs) to our operations, many of which are single-source suppliers, including the CMO that produces TEPEZZA drug substance and the CMO that produces all of our KRYSTEXXA drug substance in Israel, which is affected by the current conflict in the Middle East. Our manufacturing and supply operations may be affected by:

- capacity of manufacturing facilities;
- contamination by microorganisms or viruses, or foreign particles from the manufacturing process;
- natural or other disasters, including hurricanes, earthquakes, volcanoes or fires;
- labor disputes or shortages, including the effects of health emergencies (such as novel viruses or pandemics) or natural disasters;

- compliance with regulatory requirements;
- changes in forecasts of future demand;
- timing and actual number of production runs and production success rates and yields;
- updates of manufacturing specifications;
- contractual disputes with our suppliers and contract manufacturers;
- timing and outcome of product quality testing;
- power failures and/or other utility failures;
- cyberattacks on supplier systems;
- breakdown, failure, substandard performance or improper installation or operation of equipment (including our information technology systems and network-connected control systems or those of our contract manufacturers or third-party service providers);
- delays in the ability of the FDA or foreign regulatory agencies to provide us necessary reviews, inspections and approvals, including as a result of a subsequent extended U.S. federal or other government shutdowns;
- tariffs or other trade barriers that increase costs, limit availability, or disrupt the flow of goods; and/or
- geopolitical conflicts (such as the ongoing conflicts in Ukraine and the Middle East).

If any of these or other problems affect production in one or more of our facilities or those of our third-party contract manufacturers, or if we do not accurately forecast demand for our products or the amount of our product candidates required in clinical trials, we may be unable to start or increase production in our unaffected facilities to meet demand, particularly where such facilities are operating at or near capacity. If the efficient manufacture and supply of our products or product candidates is interrupted, we may experience delayed shipments, delays in our clinical trials, supply constraints, stock-outs, adverse event trends, contract disputes and/or recalls of our products. From time to time, we have initiated recalls of certain lots of our products. For example, in July 2014 we initiated a voluntary recall of an Aranesp lot distributed in the EU after particles were detected in a quality control sample following distribution of that lot, and in April 2018 we initiated a precautionary recall of two batches of Vectibix distributed in Switzerland after potential crimping defects were discovered in the metal seals on some product vials. If we are at any time unable to provide an uninterrupted supply of our products to patients, we may lose patients and physicians may elect to prescribe competing therapeutics instead of our products, which could have a material adverse effect on our product sales, business and results of operations.

We are expanding our manufacturing capabilities to support current and anticipated demand for our products and product candidates. These expansion efforts are complex, capital-intensive and subject to significant risks, including construction, validation and quality assurance challenges; delays or unfavorable outcomes from regulatory inspections or approvals; labor availability and workforce readiness issues; supply chain disruptions; and contractual disputes with suppliers or contractors. We may be unable to realize anticipated efficiencies or returns on our investments if there are delays or failures of our product development programs or our receipt of regulatory approvals, if our new or expanded facilities require extended periods to achieve planned production capacity or expected yields, or if demand for our products or product candidates changes or fails to materialize. These efforts often rely on a single or limited number of vendors and suppliers, and identifying or qualifying alternatives may not be feasible or may require significant time and expense due to the specialized nature of our manufacturing requirements. In addition, tariffs on imported equipment, construction materials or key inputs have increased our costs to a limited extent in 2025, and, going forward, such tariffs, or other tariffs imposed in the future, could further increase costs, potentially disrupt supply chains, and put at risk the timely and cost-effective execution of these projects.

Our manufacturing processes, those of our third-party contract manufacturers and those of certain of our third-party service providers must undergo regulatory approval processes and are subject to continued review by the FDA and other regulatory authorities. It can take longer than five years to build, validate and license another manufacturing plant, and it can take longer than three years to qualify and license a new contract manufacturer or service provider. If we elect or are required to make changes to our manufacturing processes because of new regulatory requirements, new interpretations of existing requirements or other reasons, this could increase our manufacturing costs and result in delayed shipments, delays in our clinical trials, supply constraints, stock-outs, adverse event trends or contract negotiations or disputes. Such manufacturing challenges may also occur if our existing contract manufacturers are unable or unwilling to timely implement such changes, or at all.

In addition, regulatory agencies conduct routine monitoring and inspections of our manufacturing facilities and processes as well as those of our third-party contract manufacturers and service providers. If regulatory authorities determine that we or our third-party contract manufacturers or certain of our third-party service providers have violated regulations, they may mandate corrective actions and/or issue warning letters, or even restrict, suspend or revoke our prior approvals, prohibiting us from manufacturing our products or conducting clinical trials or selling our marketed products, either until we or the affected third-party contract manufacturers or third-party service providers comply, or indefinitely. See *Our current products and products in development cannot be sold without regulatory approval*. Such issues may also delay the approval of product candidates we have submitted for regulatory review, even if such product candidates are not directly related to the products, devices or processes at issue with regulators. Regulatory authorities may also interpret issues identified with respect to a single manufacturing line as reflecting broader quality or procedural concerns that apply to other lines or facilities, which could delay approvals or adversely affect operations at other facilities. Because our third-party contract manufacturers and certain of our third-party service providers are subject to the FDA and foreign regulatory authorities, alternative qualified third-party contract manufacturers and third-party service providers may not be available on a timely basis, or at all. If we or our third-party contract manufacturers or third-party service providers cease or interrupt production or if our third-party contract manufacturers and third-party service providers fail to supply materials, products or services to us, we may experience delayed shipments, delays in our clinical trials, supply constraints, contract disputes, stock-outs and/or recalls of our products. Additionally, we distribute a substantial volume of our commercial products through our primary distribution centers in Louisville, Kentucky for the United States and in Breda, Netherlands for Europe and much of the rest of the world. We also conduct most of the labeling and packaging of our products distributed in Europe and much of the rest of the world in Breda. Our ability to timely supply products is dependent on the uninterrupted and efficient operations of our distribution and logistics centers, our third-party logistics providers and our labeling and packaging facility in Breda. Further, we rely on commercial transportation, including air and sea freight, for the distribution of our products to our customers, which has been negatively affected by pandemics, labor unrest, natural disasters and geopolitical security threats.

Changes in laws or regulations with respect to the use and/or presence of certain chemicals in our products or the components used in the research, development, manufacture and/or packaging of our products could also disrupt or restrict our ability to develop, produce or sell our products. For example, the EU, Canada, the U.S. Congress, the U.S. Environmental Protection Agency, and several U.S. states are considering legislation and/or policies to address the reporting, presence, and/or use, of certain chemicals in certain of the components used in the manufacture or packaging of commercial products, including chemicals known as per- and polyfluorinated substances (PFAS). In 2024, Canada (through Environment and Climate Change Canada) issued a notice requiring reporting on PFAS manufacture, import, and use in Canada. In addition, proposed legislation in several jurisdictions are under consideration to prohibit or otherwise regulate the importation, manufacture, or distribution of goods containing PFAS, and some such proposals do not provide exemptions for drug products, medical devices, their packaging, or the materials used in the research, development, or manufacture of such products or devices. For example, the EU is considering a ban on PFAS in the manufacturing and packaging of pharmaceutical products that could affect pharmaceutical research and development activities and commercial distribution. Some proposals, if enacted without exemptions for pharmaceutical products, and materials used in their research, development, and manufacture, or without adequate time to research and develop or otherwise identify alternative materials or suppliers, may cause significant disruptions to our ability to manufacture and supply products to the affected jurisdictions, potentially resulting in a material adverse effect on our business.

There have also been legislative and administrative proposals seeking to incentivize greater drug manufacturing in the United States with the stated goal of improving supply reliability in the United States. For example, one legislative proposal would have prohibited the U.S. Department of Veterans Affairs from purchasing certain drugs that have active pharmaceutical ingredients manufactured outside the United States. While we perform a substantial majority of our commercial manufacturing activities in the United States, including in the U.S. territory of Puerto Rico, and a substantial majority of our clinical manufacturing activities at our facility in Thousand Oaks, California, the passage of such legislation could result in foreign governments enacting retaliatory legislation or regulatory actions, which may have an adverse effect on our product sales, business and results of operations.

Our business and operations may be negatively affected by the failure, or perceived failure, of achieving our sustainability objectives.

We continue to work towards operating our business in a sustainable manner. Stakeholders, including our investors and our employees, have increasingly focused on, and are expected to continue to focus on, our sustainability practices. Policymakers, regulators and investors globally have increased their focus on sustainability matters, resulting in rapidly evolving and diverging expectations and standards. For example, California recently enacted the Climate Corporate Data Accountability Act that requires, among other things, disclosure of greenhouse gas emissions. In contrast, a number of states have adopted or proposed laws or policies that restrict the consideration of sustainability or other non-financial factors in connection with certain investment or business decisions, which may conflict with the expectations of some of our stakeholders. If our sustainability practices fail to meet stakeholders' expectations and applicable standards, or if we fail to comply with

sustainability-related regulations across our global business, there could be a material adverse effect on our reputation, business and, ultimately, our stock price.

Our sustainability report is made available on our website and describes our current sustainability goals and the progress we have made on the sustainability issues that we believe our external and internal stakeholders consider to be important, based on surveys, interviews and certain frameworks for corporate responsibility. Achieving our sustainability goals requires long-term investments and broad, coordinated activity, and we may be required to incur additional costs or allocate additional resources towards monitoring, reporting and implementing our sustainability programs. Further, we may fail to accurately assess our stakeholders' sustainability priorities and concerns, as such priorities and concerns have been rapidly changing. While we have achieved most of our goals set in prior years, whether we can achieve our current and future sustainability goals continues to be uncertain and remains subject to numerous risks, including evolving regulatory requirements and social expectations affecting sustainability practices, our ability to recruit, develop and retain a diverse workforce, the availability of suppliers and collaboration partners that can meet our environmental goals, the effects of the organic growth of our business and potential acquisitions of other businesses on our sustainability performance, and the availability and cost of technologies or resources, such as carbon credits, that support our goals. Any failure or perceived failure to meet our sustainability program priorities could result in a material adverse effect on our reputation, business and stock price.

The effects of global climate change and related natural disasters could negatively affect our business and operations.

Many of our operations and facilities, including those essential to our manufacturing, R&D and distribution activities, are in locations that are subject to natural disasters, including droughts, fires, extreme temperatures, hurricanes, tropical storms and/or floods. For example, in 2017 Hurricane Maria caused catastrophic damage, compounded in 2022 by Hurricane Fiona, to the U.S. territory of Puerto Rico, where we perform a substantial majority of our commercial manufacturing activities. Although our site was well-protected and suffered minimal damage, there can be no assurances that we would have similar results in the face of future natural disasters. The severity and frequency of weather-related natural disasters has been amplified, and is expected to continue to be amplified by, global climate change. For example, in January 2025, Los Angeles county experienced unprecedented wildfires, and while the natural disaster did not impact our facilities or their operations, a number of our staff members lost their homes or were subject to evacuation orders and/or multiple-day power outages. Such natural disasters have caused, and in the future may cause, damage to and/or disrupt our operations, which may result in a material adverse effect on our product sales, business and results of operations. Our suppliers, vendors and business partners also face similar risks, and any disruption to their operations could have an adverse effect on our supply and manufacturing chain. Further, many of our key facilities are located on islands, including Puerto Rico, Singapore and Ireland, which rely on essential port facilities that may be vulnerable to climate change-related or other natural disasters. Although we have detailed business continuity plans in place and periodic assessments of our natural disaster risk, any natural disaster may also result in prolonged interruption to our critical operational and business activities, and we may be required to incur significant costs to remedy the effects of such natural disasters and fully resume operations, which may result in a material adverse effect on our product sales, business and results of operations. See *We perform a substantial majority of our commercial manufacturing activities at our facility in the U.S. territory of Puerto Rico and a substantial majority of our clinical manufacturing activities at our facility in Thousand Oaks, California; significant disruptions or production failures at these facilities could significantly impair our ability to supply our products or continue our clinical trials and Manufacturing difficulties, disruptions or delays could limit supply of our products and limit our product sales.*

GENERAL RISK FACTORS

Global economic conditions may negatively affect us and may magnify certain risks that affect our business.

Our operations and performance have been affected, and may continue to be affected, by uncertain global economic conditions, including those arising from geopolitical and trade policy tensions and market volatility. In addition, fiscal and budgetary pressures in the United States and other jurisdictions, including uncertainty around, or reprioritization of, government funding and constrained government resources, may disrupt government operations and regulatory activities and increase pressure on healthcare budgets and reimbursement policies. See *Our sales depend on coverage and reimbursement from government and commercial third-party payers, and pricing and reimbursement pressures have affected, and are likely to continue to affect, our profitability.* As a result of global economic conditions, some third-party payers may delay or be unable to satisfy their reimbursement obligations. Job losses or other economic hardships (including inflation) may also affect patients' ability to afford healthcare as a result of increased co-pay or deductible obligations, greater cost sensitivity to existing co-pay or deductible obligations, lost healthcare insurance coverage or for other reasons. We believe such conditions have led and could continue to lead to reduced demand for our products, which could have a material adverse effect on our product sales, business and results of operations. Our operational costs, including the cost of energy, materials, labor, distribution and our other operational and facilities costs are subject to market conditions and have been adversely affected by tariffs to a limited extent. Although we monitor our distributors', customers' and suppliers' financial condition and their liquidity to mitigate our business risks, some of our distributors, customers and suppliers may become insolvent, which could have a material adverse effect on

our product sales, business and results of operations. A significant worsening of global economic conditions could precipitate or materially amplify the other risks described herein. On April 2, 2025, the Administration issued an executive order (the April 2025 Tariff EO) imposing a universal 10% tariff on all imported goods, with certain exceptions including pharmaceuticals. The April 2025 Tariff EO imposed additional higher tariffs on approximately 60 countries with which the United States has trade deficits. Recent and ongoing changes in U.S. trade and tariff policies, including the imposition, modification, suspension and threatened expansion of tariffs on imported goods, as well as retaliatory measures by foreign governments, have increased uncertainty in the overall business and operating environment. Although certain pharmaceutical products are exempt from some tariffs, these measures have increased, and may continue to increase, our costs, including for materials, components and supplies used in research and development and manufacturing activities, have adversely affected, and may continue to adversely affect, our operations in various markets, and could disrupt global supply chains. On October 24, 2025, the Administration initiated, under the Trade Act of 1974, a Section 301 investigation of China's implementation of the Economic and Trade Agreement between the U.S. and Chinese governments. This investigation, and any other 301 investigations initiated, may result in additional tariffs on imported goods from China and any other foreign markets subsequently investigated, respectively, potentially including pharmaceutical products and other goods that Amgen requires for the manufacture of our products. If subject to Section 301 tariffs, China, and other affected foreign governments, may retaliate against such tariffs by imposing tariffs of their own on U.S.-made goods. Given the many uncertainties and variables, it is currently unclear the extent, and degree, to which existing and future tariffs will disrupt and adversely affect our business activities (including product sales, and conduct of clinical trial and research and development activities), and the global economic environment, and/or amplify the other risks described herein.

We maintain a significant portfolio of investments on our consolidated balance sheets. Global capital markets have experienced, and may continue to experience, periods of volatility and disruption, including as a result of interest rate fluctuations, inflation, liquidity conditions, credit market stress and geopolitical events. Certain of our assets, including equity investments, are exposed to market fluctuations that, in a sustained or recurrent series of market disruptions, could result in impairments or losses on sale. The value of our investments may also be adversely affected by interest rate fluctuations, inflation, downgrades in credit ratings, illiquidity in the capital markets, geopolitical events and other factors that may result in other-than-temporary declines in the value of our investments. Any of those events could cause us to record impairment charges with respect to our investment portfolio or to realize losses on sales of investments. We also maintain a majority of our cash and cash equivalents in accounts with major multi-national financial institutions, and our deposits at these institutions exceed insured limits. Market conditions can adversely affect the viability of these institutions. In the event of failure of any of the financial institutions where we maintain our cash and cash equivalents, there can be no assurance that we would be able to access uninsured funds in a timely manner or at all. Inability to access, or a delay in accessing these funds, could adversely affect our business and financial position.

Our stock price is volatile.

Our stock price, like that of our peers in the biotechnology and pharmaceutical industries, is volatile. Our revenues and operating results may fluctuate from period to period for a number of reasons. Events such as a delay in product development, changes to our expectations or strategy or even a relatively small revenue shortfall may cause financial results for a period to be below our expectations or projections. As a result, our revenues and operating results and, in turn, our stock price may be subject to significant fluctuations. Announcements or discussions, including via social media channels, of possible restrictive actions by government or private payers that would negatively affect our business or industry if ultimately enacted or adopted may also cause our stock price to fluctuate, whether or not such restrictive actions ever actually occur. Similarly, actual or perceived safety issues with our products or similar products or unexpected clinical trial results can have an immediate and rapid effect on our stock price, whether or not our operating results are materially affected.

Item 1B. UNRESOLVED STAFF COMMENTS

None.

Item 1C. CYBERSECURITY

Risk Management and Strategy

Amgen has a multi-layered and iterative approach towards assessing, identifying, managing and mitigating risks from cybersecurity threats. The Amgen Technology & Medical Organizations (ATMOS) function is designed to support our productivity, innovation and outreach globally through the quality delivery of information systems, solutions and services for our business and operations. The ATMOS function has a Cybersecurity & Digital Trust (CDT) team that assesses and reduces cybersecurity exposure, including by providing employees with training and resources to identify potential cybersecurity threats and implementing information technology security practices. The CDT team also monitors for cybersecurity threat activity and seeks to mitigate the impact from cybersecurity incidents by deploying information security engineers, system architects, analysts and cybersecurity specialists to provide monitoring, reporting and management of cybersecurity incidents.

To evaluate the progress of its activities, our ATMOS function uses various industry and regulatory frameworks as guides to assess the state of the Company's cybersecurity program maturity and controls, including our organizational, people, physical and technological controls. The CDT team also conducts reviews and evaluations of our cybersecurity resilience program with Amgen's Cybersecurity & Digital Trust Governance Council (which includes leaders from CDT, Worldwide Compliance and Business Ethics, Regulatory Affairs, Operations, R&D, Global Commercial Operations, Corporate Audit, Finance, Law and Business Development functions).

Our cybersecurity risk management program is considered by and integrated into our Company-wide Enterprise Risk Management program and shares common methodologies, reporting channels and governance processes that apply across the Enterprise Risk Management program to that of other enterprise level risks (such as product development, safety and surveillance, financial and intellectual property risks). Regular evaluations are conducted of the greatest risks to our business and their underlying risk drivers as well as the associated mitigation activities, maturity and controls. This program is overseen by our Executive Vice President and Chief Financial Officer and guided by the Enterprise Risk Council, a cross-functional group of the Company's business leaders representing key business functions that is co-chaired by our Chief Audit Executive. The results of the enterprise risk evaluations and the status and operation of the Enterprise Risk Management program are presented to our Board of Directors, which oversees the Company's enterprise-level risks.

Further, our corporate audit function is responsible for assessing risk and testing whether, and the extent to which, our information security policies and practices are being implemented effectively within our business and by third party providers. Findings from such reports and related corrective action plans are shared with our CDT team, Company leadership, and the Audit Committee and Corporate Responsibility and Compliance Committee (CRCC) of our Board of Directors.

In addition to leveraging the Company's own information technology resources, our Incident Response and Cyber Threat Intelligence teams engage, as needed, third-party cybersecurity risk assessors and consultants to assist in recognizing threats, identifying security vulnerabilities and evaluating the impact of cybersecurity attacks and incidents when they occur. On a biennial basis, our ATMOS also engages external third-party experts to assess the Company's cybersecurity control maturity across the organization and develops plans to address such experts' recommendations.

Our CDT function has processes to oversee and identify the risks of cybersecurity threats associated with third-party service providers and monitors and works to mitigate the impact of cybersecurity incidents encountered by our third-party service providers. Upon becoming aware of cybersecurity incidents encountered by our third-party service providers, the CDT function's Incident Response and Cyber Threat Intelligence teams are deployed to evaluate and mitigate the impact of such incidents on our business.

In connection with our adoption of artificial intelligence (AI) tools in our business, including AI tools customized for our business and a variety of Amgen-built tools for use across applications, the Company established an AI Governance Council composed of cross-functional leadership that oversees the safe adoption of third-party AI services, including by establishing guardrails to reduce risks and allocating resources to provide staff training on the proper use of AI and responsible AI practices. The AI Governance Council is co-sponsored by our Chief Compliance Officer and Senior Vice President, Artificial Intelligence & Data.

Despite our layered controls and cybersecurity efforts, the Company and its third-party vendors have experienced cyberattacks and information security vulnerabilities, and while such incidents have not had a material adverse effect on the Company, there can be no assurance that future cybersecurity attacks or incidents would not result in a material adverse effect

on our business strategy, results of operations or financial condition. For examples of such matters and a discussion of the risks that we face, see Item 1A. Risk Factors—*A breakdown of our information technology systems, cyberattack or information security breach could significantly compromise the confidentiality, integrity and availability of our information technology systems, network-connected control systems and/or our data, interrupt the operation of our business and/or affect our reputation.* However, we have not identified risk from known cybersecurity threats, including as a result of any prior cybersecurity incidents, that have materially affected or are reasonably likely to materially affect us, including our operations, business strategy, results of operations or financial condition.

Governance

Our Board of Directors oversees an enterprise-wide approach to risk management, including risks related to information systems and cybersecurity, and each Board committee has primary risk oversight responsibilities aligned with its areas of focus. At each regular meeting of the Board, the Board receives and considers reports from each of its committees, and such reports provide additional detail on significant risk management issues as appropriate, including cybersecurity. The CRCC is the committee that has primary oversight responsibility for the Company's information systems and management of cybersecurity and receives reports from our Senior Vice President and Chief Information Officer (CIO) and Chief Information Security Officer (CISO) that includes reviews of our information systems strategy, technology investments, cybersecurity risks and incidents, and third-party risk management, as well as an annual evaluation of the Company's cybersecurity status. The Board's Audit Committee has oversight responsibility of our internal controls, assurances and financial risks. The Audit Committee is provided with copies of materials presented to our CRCC by our CIO and CISO and receives reports from our CIO regarding topics including integration or implementation of new financial systems and key controls and governance designed to address cybersecurity risks associated with the use of such new financial systems.

Our management team, including our CIO and CISO, supervises efforts to prevent, detect, mitigate and remediate cybersecurity risks and incidents through various means, which may include briefings from internal information security personnel; threat intelligence and other information obtained from governmental, public or private sources, including external consultants engaged by us; and alerts and reports produced by security tools deployed in the information systems environment.

Our CISO, who heads our CDT team and is accountable for the Company's cybersecurity risk management program, joined the Company's information systems organization in 2016, is a Certified Information Systems Security Professional and is certified in risk and information systems control. Previously, our CISO served in both leadership and operational positions as a cybersecurity professional in the U.S. government and was a cybersecurity consultant, providing a wide range of cybersecurity services to various U.S. government agencies and departments. Our CISO is overseen by our CIO, who has 28 years of experience in information systems (including over 15 years at the Company and more than 6 years as a senior technology executive outside of Amgen), and holds a Computer Information Systems B.S. and an Information Technology Management MBA. Our Executive Vice President and Chief Technology Officer (CTO) leads our ATMOS function and oversees our CIO.

As leaders of the Technology and CDT functions within ATMOS, respectively, the Company's CIO and CISO are informed about and monitor significant cybersecurity threats and incidents through the Company's internal cybersecurity reporting structure. Our CDT team is responsible for monitoring and detecting cybersecurity threats and incidents. Our CDT team, overseen by our CISO, is also responsible for the mitigation and remediation of cybersecurity incidents. When members of the CDT team detect a cybersecurity threat or incident or are made aware of a cybersecurity incident encountered by a third-party service provider, the discovery is communicated to the Incident Response team, which includes our CISO and other senior members of the CDT function. The Incident Response team evaluates the severity of the cybersecurity threat or incident and shares its findings with our CISO.

Our CISO and/or his senior team leaders, in addition to our CIO and CTO, also provide regular reports to executives leading our Finance, Compliance, Law and Human Resources functions on potentially significant cybersecurity incidents and the progress made towards mitigation and remediation of those incidents. These leaders oversee reporting to our CRCC and Audit Committee, and reporting of such cybersecurity incidents is included in the course of regular meetings of such committees. Additionally, in appropriate circumstances, reporting of potentially significant cybersecurity incidents is made directly to the leaders of our CRCC and Audit Committee or directly to the Board of Directors outside of their regular meeting schedule. Further, in support of our internal controls, our CISO also reviews cybersecurity matters and trends with our Accounting and Law functions at least on a quarterly basis.

In 2025, we substantially transitioned or decommissioned the technology systems from the legacy Horizon environment to the Amgen environment. The remaining systems are being monitored by the Amgen cybersecurity team in accordance with the governance processes detailed above.

Item 2. PROPERTIES

As of December 31, 2025, we owned or leased approximately 160 properties. The locations and primary functions of significant properties are summarized in the following tables:

U.S. Location:	Manufacturing	Administrative	R&D	Sales and marketing	Warehouse	Distribution center
Thousand Oaks, CA ⁽¹⁾	x	x	x	x	x	x
Juncos, Puerto Rico	x	x			x	x
West Greenwich, RI	x	x			x	
Deerfield, IL		x	x	x		
Cambridge, MA			x			
New Albany, OH	x	x			x	
San Francisco, CA			x			
Tampa, FL		x		x		
Louisville, KY					x	x
Other U.S. cities ⁽²⁾		x	x	x		

ROW Location:	Manufacturing	Administrative	R&D	Sales and marketing	Warehouse	Distribution center
Canada		x	x	x		
China		x		x		
Denmark		x	x	x		
Germany		x	x	x		
Iceland		x	x			
India		x				
Ireland	x	x		x	x	x
Netherlands	x	x		x	x	x
Singapore	x	x		x	x	
Other countries ⁽²⁾		x	x	x	x	

⁽¹⁾ Corporate headquarters.

⁽²⁾ Includes smaller properties in other U.S. and ROW locations, primarily for administrative and sales and marketing.

Excluded from the information above are (i) undeveloped land and leased properties that have been abandoned and (ii) certain buildings we still own but that are no longer used in our business. Additionally, in 2025 we broke ground on our second drug substance manufacturing facility in Holly Springs, North Carolina; announced expansions of our manufacturing network in Ohio and the U.S. territory of Puerto Rico; and broke ground on a new, state-of-the-art science and innovation center at our corporate headquarters in Thousand Oaks, California. There are no material encumbrances on our owned properties.

We believe our facilities are suitable for their intended uses and, in conjunction with our third-party contract manufacturing agreements, provide adequate capacity and are sufficient to meet our expected needs. See Item 1A. Risk Factors for a discussion of the factors that could adversely impact our manufacturing operations and the global supply of our products.

See Item 1. Business—Manufacturing, Distribution and Raw Materials.

Item 3. LEGAL PROCEEDINGS

Certain of the legal proceedings in which we are involved are discussed in Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements and are hereby incorporated by reference.

Item 4. MINE SAFETY DISCLOSURES

Not applicable.

PART II

Item 5. MARKET FOR REGISTRANT'S COMMON EQUITY, RELATED STOCKHOLDER MATTERS AND ISSUER PURCHASES OF EQUITY SECURITIES

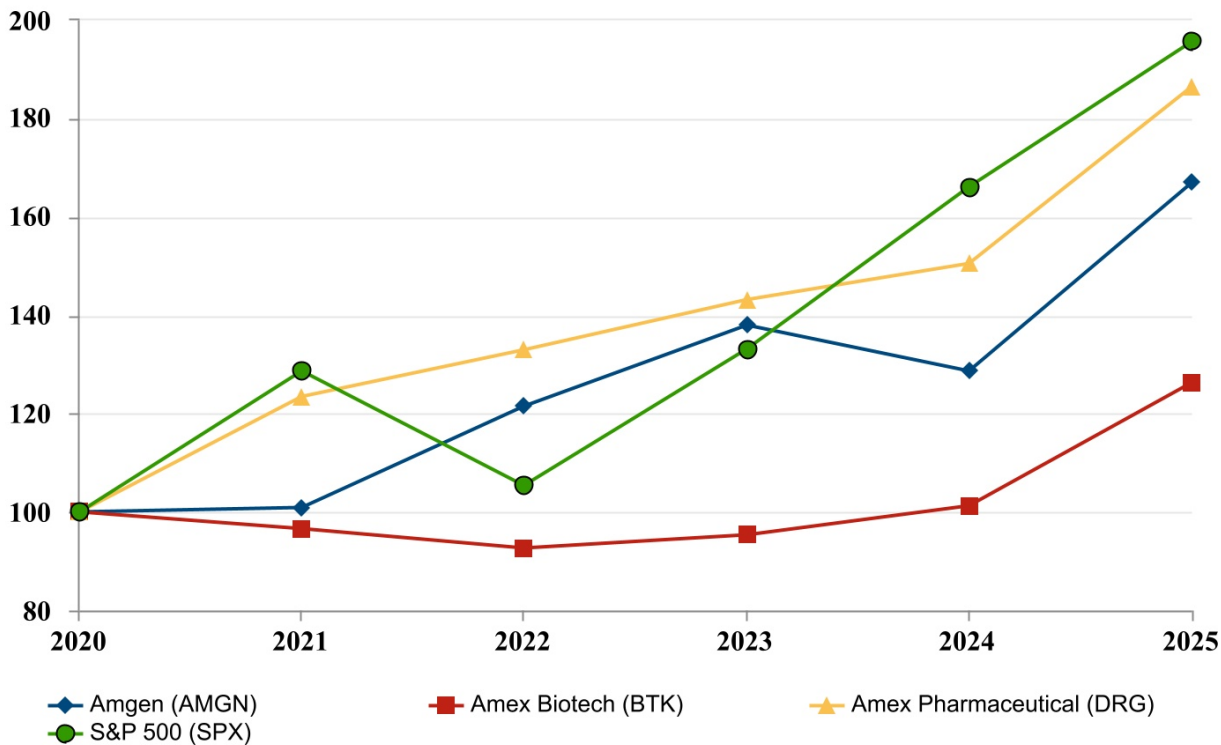
Common stock

Our common stock trades on the Nasdaq Global Select Market under the symbol AMGN. As of February 10, 2026, there were approximately 3,824 holders of record of our common stock.

Performance graph

The following graph shows the value of an investment of \$100 on December 31, 2020, in each of Amgen common stock, the Amex Biotech Index, the Amex Pharmaceutical Index and Standard & Poor's 500 Index. All values assume reinvestment of the pretax value of dividends and are calculated as of December 31 of each year. The historical stock price performance of the Company's common stock shown in the performance graph is not necessarily indicative of future stock price performance.

**Comparison of Five-Year Cumulative Total Return
of a \$100 Investment on December 31, 2020**



	12/31/2020	12/31/2021	12/31/2022	12/31/2023	12/31/2024	12/31/2025
Amgen (AMGN)	\$100.00	\$100.87	\$121.54	\$137.97	\$128.71	\$166.88
Amex Biotech (BTK)	\$100.00	\$96.48	\$92.61	\$95.27	\$101.21	\$126.28
Amex Pharmaceutical (DRG)	\$100.00	\$123.37	\$132.94	\$143.21	\$150.48	\$186.40
Standard & Poor's 500 (SPX)	\$100.00	\$128.68	\$105.29	\$133.03	\$165.98	\$195.62

The material in the above performance graph is not soliciting material, is not deemed filed with the SEC and is not incorporated by reference in any filing of the Company under the Securities Act or the Exchange Act, whether made on, before or after the date of this filing and irrespective of any general incorporation language in such filing.

Stock repurchase program

During the year ended December 31, 2025, we had one outstanding stock repurchase program, under which repurchase activity was as follows:

	Total number of shares purchased	Average price paid per share	Total number of shares purchased as part of publicly announced program	Maximum dollar value that may yet be purchased under the program
October 1–October 31	—	\$ —	—	\$ 6,779,253,902
November 1–November 30	—	\$ —	—	\$ 6,779,253,902
December 1–December 31	—	\$ —	—	\$ 6,779,253,902
	—		—	
January 1–December 31 ⁽¹⁾	—	\$ —	—	

⁽¹⁾ During the year ended December 31, 2025, the Company purchased 1,700 shares at an average price paid of \$284.67 per share from staff members to satisfy federal law compliance obligations. These shares were not repurchased under our stock repurchase program.

Dividends

For the years ended December 31, 2025 and 2024, we paid quarterly dividends. We expect to continue to pay quarterly dividends, although the amount and timing of any future dividends are subject to approval by our Board of Directors. Additional information required by this item is incorporated herein by reference to Part IV—Note 17, Stockholders’ equity, to the Consolidated Financial Statements.

Securities Authorized for Issuance Under Existing Equity Compensation Plans

Information about securities authorized for issuance under existing equity compensation plans is incorporated by reference from Part III, Item 12—Securities Authorized for Issuance Under Existing Equity Compensation Plans.

Item 6. RESERVED

Item 7. MANAGEMENT’S DISCUSSION AND ANALYSIS OF FINANCIAL CONDITION AND RESULTS OF OPERATIONS

The following MD&A is intended to assist the reader in understanding Amgen’s business. MD&A is provided as a supplement to, and should be read in conjunction with, our consolidated financial statements and accompanying notes. Our results of operations discussed in MD&A are presented in conformity with GAAP. Amgen operates in one operating segment: human therapeutics. Therefore, our results of operations are discussed on a consolidated basis.

Forward-looking statements

This report and other documents we file with the SEC contain forward-looking statements that are based on current expectations, estimates, forecasts and projections about us, our future performance, our business, our beliefs and our management’s assumptions. In addition, we, or others on our behalf, may make forward-looking statements in press releases, written statements or our communications and discussions with investors and analysts in the normal course of business through meetings, webcasts, phone calls and conference calls. Such words as “expect,” “anticipate,” “outlook,” “could,” “target,” “project,” “intend,” “plan,” “believe,” “seek,” “estimate,” “should,” “may,” “assume” and “continue” as well as variations of such words and similar expressions are intended to identify such forward-looking statements. These statements are not guarantees of future performance and they involve certain risks, uncertainties and assumptions that are difficult to predict. We describe our respective risks, uncertainties and assumptions that could affect the outcome or results of operations in Part I, Item 1A. Risk Factors. We have based our forward-looking statements on our management’s beliefs and assumptions based on information available to our management at the time the statements are made. We caution you that actual outcomes and results may differ materially from what is expressed, implied or forecasted by our forward-looking statements. Reference is made in particular to forward-looking statements regarding product sales, regulatory activities, clinical trial results, reimbursement, expenses, EPS, liquidity and capital resources, trends, planned dividends, stock repurchases and collaborations. Except as required under the federal securities laws and the rules and regulations of the SEC, we do not have any intention or obligation to update publicly any forward-looking statements after the distribution of this report, whether as a result of new information, future events, changes in assumptions or otherwise.

Overview

Amgen Inc. (including its subsidiaries, referred to as “Amgen,” “the Company,” “we,” “our” or “us”) discovers, develops, manufactures and delivers innovative medicines to fight some of the world’s toughest diseases. We focus on areas of high unmet medical need and leverage our expertise to strive for solutions that dramatically improve people’s lives, while also reducing the social and economic burden of disease. We helped launch the biotechnology industry more than 45 years ago and have grown to be one of the world’s leading independent biotechnology companies. Our robust pipeline includes potential first-in-class medicines at all stages of development.

Our principal products are Prolia, Repatha, Otezla, ENBREL, EVENITY, XGEVA, TEPEZZA, BLINCYTO, Nplate, TEZSPIRE, KYPROLIS, Aranesp, KRYSTEXXA and Vectibix. We also market a number of other products, including but not limited to MVASI, PAVBLU, UPLIZNA, IMDELLTRA/IMDYLLTRA, AMJEVITA/AMGEVITA, TAVNEOS, Neulasta, LUMAKRAS/LUMYKRAS, RAVICTI, Parsabiv, Aimovig, WEZLANA/WEZENLA and PROCYSBI. For additional information about our products, see Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products.

Our strategy is the integrated set of actions we take to improve our competitive position in the industry. In 2025, we generated strong sales growth across our product portfolio and regions; advanced our innovative pipeline; and continued to expand and enhance our world-class manufacturing network. We accomplished these objectives while maintaining a strategic and disciplined approach to capital allocation, including retiring \$6.0 billion of debt.

In 2025, we achieved several significant regulatory, clinical and operational milestones. We obtained multiple regulatory approvals, including new indications for UPLIZNA and TEZSPIRE; a broadened FDA approval for Repatha; and full FDA approval for IMDELLTRA for the treatment of ES-SCLC. We also advanced our innovative pipeline, including the initiation of six global Phase 3 clinical studies for MariTide and the reporting of Phase 3 data across several programs. In addition, we continued to invest in expanding and enhancing our manufacturing capacity, including facilities in Ohio, North Carolina and the U.S. territory of Puerto Rico. Furthermore, in 2025 we also broke ground on a new state-of-the-art R&D facility in Thousand Oaks, California, to further enhance collaboration and innovation across R&D and process development activities. For additional information on our pipeline and clinical development updates, see Part I, Item 1. Business—Research and Development and Selected Product Candidates, and Part I, Item 1. Business—Significant Developments. For additional information on our manufacturing operations, see Part I, Item 1. Business—Manufacturing, Distribution and Raw Materials.

Total product sales increased 10% in 2025, driven by volume growth of 13%, partially offset by declines in net selling price of 3%.

Cash flows from operating activities in 2025 totaled \$10.0 billion, which supported investment in our business, including capital expenditures of \$1.9 billion to enhance and expand our manufacturing network, and allowed us to both reduce our debt and return capital to shareholders through the payment of cash dividends. For 2025, we retired \$6.0 billion of debt and increased our quarterly cash dividend by 6% to \$2.38 per share of common stock. In December 2025, the Board of Directors declared a cash dividend of \$2.52 per share of common stock for the first quarter of 2026, an increase of 6% over the same period in the prior year, to be paid in March 2026.

Amgen's approach to human capital management focuses on attracting, developing and retaining a highly skilled global workforce to support the discovery, development and commercialization of innovative medicines. Our compensation, benefits and development programs are designed to promote performance, accountability, adherence to Company values and alignment with shareholder interests. We believe our culture supports innovation, collaboration and productivity as we execute on our mission to serve patients. For additional information, see Part I, Item 1. Business—Human Capital Resources.

We have a long-standing ambition to be environmentally responsible, and we regularly set targets to challenge ourselves to deliver further improvements. As part of our environmental sustainability efforts, we have established long-term targets to meet by 2027, including achieving carbon neutrality, reducing water consumption by 40% and reducing waste disposed by 75%.^{2,3}

Our long-term success depends, to a great extent, on our ability to continue to discover, develop and commercialize innovative products and acquire or collaborate on therapies currently in development by other companies. We must grow sales from existing and new products to achieve revenue growth and to offset revenue losses caused by products' loss of their exclusivity or launches of competing products. For example, our patents for RANKL antibodies, including sequences, for Prolia and XGEVA expired in February 2025 in the United States and in November 2025 in select countries in Europe. Certain of our products face increasing pressure from competition, including biosimilars and generics. For additional information, including information on the expirations of patents for various products, see Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Patents*, and Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Competition*. We devote considerable resources to R&D activities, but successful product development in the biotechnology industry is highly uncertain. We also face increasing regulatory scrutiny of safety and efficacy both before and after products launch.

Tariffs and trade protection measures

Recent and ongoing changes in U.S. trade and tariff policies, including the imposition, modification, suspension and threatened expansion of tariffs on imported goods, as well as retaliatory measures by foreign governments, have increased uncertainty in the overall business and operating environment. Numerous tariffs and trade protection measures have been proposed, and in a number of cases, implemented by the United States and other countries, including the April 2025 Tariff EO, which imposed a universal 10% tariff on goods imported into the United States, with certain exceptions including pharmaceuticals. Further, there were previous proposals for sector-specific tariffs on our industry, and in December 2025, in recognition of our capital investments in U.S. manufacturing, we received relief from Section 232 tariffs, pending final determination under such section of the Trade Expansion Act of 1962, for approximately the next three years. Tariffs and trade protection measures may adversely affect our business and results of operations. For additional discussion of these and other risks, see Part I, Item 1A. Risk Factors, of this Annual Report on Form 10-K.

Macroeconomic and other challenges

Uncertain macroeconomic conditions, including the risk of inflation, fluctuating interest rates and instability in the financial system, as well as rising healthcare costs, continue to pose challenges to our business. Uncertainty around tariffs and trade protection measures in the United States and other countries, including the imposition, modification, suspension and threatened expansion of tariffs on imported goods, along with ongoing geopolitical conflicts and rising geopolitical tensions, continue to create additional uncertainty in global macroeconomic conditions. Additionally, with public and private healthcare-provider focus, the industry continues to be subject to cost containment measures and significant pricing pressures, resulting in net price declines.

Moreover, provisions of the IRA, as well as the expanded utilization of the 340B Program, have negatively affected, and are likely to continue to negatively affect, our business. For example, CMS has selected ENBREL and Otezla for Medicare price setting beginning in 2026 and 2027, respectively. In addition to the IRA, other recent and proposed U.S. policy actions focus on drug pricing, including the Most-Favored-Nations Prescription Drug Pricing Executive Order (MFN EO) and the July MFN Letter that was delivered to a number of pharmaceutical companies, including Amgen. In December 2025, we announced

² Represents reductions against established baselines, taking into account only verified reduction projects and does not take into account changes associated with contraction or expansion of the Company.

³ Carbon neutrality goal refers to Scopes 1 and 2.

that we are taking actions that satisfy the components outlined in the July MFN Letter, including the Administration's MFN pricing requests. We also announced the expansion of our direct-to-patient program. While this development reflects ongoing engagement on pricing policy, the ultimate effects on our pricing, reimbursement, net sales and profitability remain uncertain in light of such evolving regulatory and policy expectations. For additional discussion of these and other risks, see Part I, Item 1A. Risk Factors, of this Annual Report on Form 10-K.

Finally, wholesale and end-user buying patterns can affect our product sales. These buying patterns can cause fluctuations in quarterly product sales, but have generally not been significant to date when comparing full-year product performance to the prior year. See Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products, and Part I, Item 1A. Risk Factors, of this Annual Report on Form 10-K for further discussion of certain factors that could impact our future product sales.

Selected financial information

The following is an overview of our results of operations (in millions, except percentages and per-share data):

	Year ended December 31, 2025	Change	Year ended December 31, 2024
Product sales:			
U.S.	\$ 25,656	10 %	\$ 23,301
ROW	9,492	9 %	8,725
Total product sales	35,148	10 %	32,026
Other revenues	1,603	15 %	1,398
Total revenues	\$ 36,751	10 %	\$ 33,424
Operating expenses	\$ 27,671	6 %	\$ 26,166
Operating income	\$ 9,080	25 %	\$ 7,258
Net income	\$ 7,711	89 %	\$ 4,090
Diluted EPS	\$ 14.23	88 %	\$ 7.56
Diluted shares	542	0 %	541

In the following discussion of changes in product sales, any reference to volume growth or decline refers to changes in the purchases of our products by healthcare providers (such as physicians or their clinics), dialysis centers, hospitals and pharmacies. In addition, any reference to increases or decreases in inventory refers to changes in inventory held by wholesaler customers and end users (such as pharmacies).

Total product sales increased 10% in 2025, driven by volume growth of 13%, partially offset by declines in net selling price of 3%. U.S. volume grew 13% and ROW volume grew 14%, driven by volume growth in certain brands, including Repatha, PAVBLU, EVENITY, IMDELLTRA/IMDYLLTRA and TEZSPIRE.

For 2026, we expect volume growth from certain brands to be partially offset by net selling price declines. Further, the first quarter of a year historically represents the lowest product sales quarter for the year, in part due to plan changes, insurance reverifications and higher co-pay expenses as U.S. patients work through deductibles, including for ENBREL and Otezla, and to a lesser extent for KRYSTEXXA, TEZSPIRE and Repatha, particularly for products acquired through pharmacy benefit programs.

Other revenues increased 15% for 2025, primarily driven by higher royalty income.

Operating expenses increased 6% for 2025, primarily driven by investments in Later-Stage Clinical Programs and Otezla intangible asset impairment charges in 2025, partially offset by lower amortization expense from acquisition-related assets, including the fair value step-up of inventory acquired from Horizon. See Part IV—Note 13, Goodwill and other intangible assets, to the Consolidated Financial Statements, for additional information related to the Otezla intangible asset impairment charges.

Uncertain macroeconomic conditions, including uncertainty around tariffs and trade protection measures, ongoing geopolitical conflicts and rising geopolitical tensions, changes in the healthcare ecosystem, and potential government policy actions, including MFN pricing or similar drug pricing reforms, have the potential to introduce variability into product sales. Furthermore, product sales continue to be impacted by actions from governments and other entities to address macroeconomic challenges, provisions of the IRA, expanded utilization of the 340B Program and growth in numbers of Medicaid enrollees and uninsured individuals. See Part I, Item 1. Business—Reimbursement, and Part I, Item 1A. Risk Factors, of this Annual Report on Form 10-K.

Results of operations

Product sales

Worldwide product sales were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Prolia	\$ 4,414	1 %	\$ 4,374	8 %	\$ 4,048
Repatha	3,016	36 %	2,222	36 %	1,635
Otezla	2,265	7 %	2,126	(3)%	2,188
ENBREL	2,226	(33)%	3,316	(10)%	3,697
EVENITY	2,100	34 %	1,563	35 %	1,160
XGEVA	2,084	(6)%	2,225	5 %	2,112
TEPEZZA ⁽¹⁾	1,903	3 %	1,851	*	448
BLINCYTO	1,559	28 %	1,216	41 %	861
Nplate	1,524	5 %	1,456	(1)%	1,477
TEZSPIRE ⁽²⁾	1,478	52 %	972	71 %	567
KYPROLIS	1,412	(6)%	1,503	7 %	1,403
Aranesp	1,389	4 %	1,342	(1)%	1,362
KRYSTEXXA ⁽¹⁾	1,340	13 %	1,185	*	272
Vectibix	1,175	12 %	1,045	6 %	984
Other products ⁽³⁾	7,263	29 %	5,630	20 %	4,696
Total product sales	<u>\$ 35,148</u>	10 %	<u>\$ 32,026</u>	19 %	<u>\$ 26,910</u>
Total U.S.	\$ 25,656	10 %	\$ 23,301	21 %	\$ 19,272
Total ROW	9,492	9 %	8,725	14 %	7,638
Total product sales	<u>\$ 35,148</u>	10 %	<u>\$ 32,026</u>	19 %	<u>\$ 26,910</u>

* Change in excess of 100%

⁽¹⁾ TEPEZZA and KRYSTEXXA were acquired from our Horizon acquisition on October 6, 2023, and include product sales in the periods after the acquisition date.

⁽²⁾ TEZSPIRE is marketed by our collaborator AstraZeneca outside the United States.

⁽³⁾ Consists of product sales of our non-principal products.

Future sales of our products will depend in part on the factors discussed in the Overview, Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Competition*, Part I, Item 1. Business—Reimbursement, Part I, Item 1A. Risk Factors, and any additional factors discussed in the individual product sections below. In addition, for a list of our products’ significant competitors, see Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products—*Competition*.

Prolia

Total Prolia sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Prolia — U.S.	\$ 2,978	3 %	\$ 2,885	6 %	\$ 2,733
Prolia — ROW	1,436	(4)%	1,489	13 %	1,315
Total Prolia	<u>\$ 4,414</u>	<u>1 %</u>	<u>\$ 4,374</u>	<u>8 %</u>	<u>\$ 4,048</u>

The increase in global Prolia sales for 2025 was primarily driven by volume growth of 2% and favorable changes to estimated sales deductions of 2%, partially offset by lower net selling price.

The increase in global Prolia sales for 2024 was driven by volume growth.

As disclosed in Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products, our patents for RANKL antibodies, including sequences, for Prolia expired in February 2025 in the United States and in November 2025 in select countries in Europe. For 2026, we expect accelerated sales erosion driven by increased competition, as multiple biosimilars have launched in the United States and ROW.

For a discussion of ongoing litigation related to Prolia, see Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements.

Repatha

Total Repatha sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Repatha — U.S.	\$ 1,663	46 %	\$ 1,139	44 %	\$ 793
Repatha — ROW	1,353	25 %	1,083	29 %	842
Total Repatha	<u>\$ 3,016</u>	<u>36 %</u>	<u>\$ 2,222</u>	<u>36 %</u>	<u>\$ 1,635</u>

The increase in global Repatha sales for 2025 was driven by volume growth. For 2026, we expect product sales for Repatha to follow the historical pattern of lower sales in the first quarter relative to subsequent quarters, as discussed above. Additionally, for 2026, we expect net selling price to decline by approximately mid-single digits.

The increase in global Repatha sales for 2024 was primarily driven by volume growth of 43%, partially offset by lower net selling price of 10%.

For a discussion of ongoing litigation related to Repatha, see Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements.

Otezla

Total Otezla sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Otezla — U.S.	\$ 1,839	8 %	\$ 1,699	(4)%	\$ 1,777
Otezla — ROW	426	0 %	427	4 %	411
Total Otezla	<u>\$ 2,265</u>	<u>7 %</u>	<u>\$ 2,126</u>	<u>(3)%</u>	<u>\$ 2,188</u>

The increase in global Otezla sales for 2025 was primarily driven by volume growth of 3% and favorable changes to estimated sales deductions of 2%. For 2026, we expect product sales for Otezla to follow the historical pattern of lower sales in the first quarter relative to subsequent quarters, as discussed above.

In January 2025, Otezla was selected by CMS for Medicare price setting that will be applicable beginning in 2027. As a result, we expect further declines in net selling price driven by Medicare price setting beginning in 2027. See Part IV—Note 13, Goodwill and other intangible assets, to the Consolidated Financial Statements for additional information related to the Otezla intangible asset impairment charges.

The decrease in global Otezla sales for 2024 was primarily driven by lower net selling price of 8%, partially offset by volume growth of 3%.

ENBREL

Total ENBREL sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
ENBREL — U.S.	\$ 2,199	(33)%	\$ 3,288	(10)%	\$ 3,650
ENBREL — Canada	27	(4)%	28	(40)%	47
Total ENBREL	<u>\$ 2,226</u>	<u>(33)%</u>	<u>\$ 3,316</u>	<u>(10)%</u>	<u>\$ 3,697</u>

The decrease in ENBREL sales for 2025 was primarily driven by lower net selling price of 36% resulting from the impact of increased 340B Program mix, U.S. Medicare Part D redesign and higher commercial discounts, partially offset by volume growth of 4%. For 2026, we expect product sales for ENBREL to follow the historical pattern of lower sales in the first quarter relative to subsequent quarters, as discussed above.

The decrease in ENBREL sales for 2024 was driven by lower net selling price.

EVENTITY

Total EVENTITY sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
EVENTITY — U.S.	\$ 1,600	41 %	\$ 1,131	40 %	\$ 809
EVENTITY — ROW	500	16 %	432	23 %	351
Total EVENTITY	<u>\$ 2,100</u>	<u>34 %</u>	<u>\$ 1,563</u>	<u>35 %</u>	<u>\$ 1,160</u>

The increases in global EVENTITY sales for 2025 and 2024 were driven by volume growth.

XGEVA

Total XGEVA sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
XGEVA — U.S.	\$ 1,355	(10)%	\$ 1,507	(1)%	\$ 1,527
XGEVA — ROW	729	2 %	718	23 %	585
Total XGEVA	<u>\$ 2,084</u>	<u>(6)%</u>	<u>\$ 2,225</u>	<u>5 %</u>	<u>\$ 2,112</u>

The decrease in global XGEVA sales for 2025 was driven by lower volume.

The increase in global XGEVA sales for 2024 was driven by higher net selling price.

As disclosed in Part I, Item 1. Business—Marketing, Distribution and Selected Marketed Products, our patents for RANKL antibodies, including sequences, for XGEVA expired in February 2025 in the United States and in November 2025 in select countries in Europe. For 2026, we expect accelerated sales erosion driven by increased competition, as multiple biosimilars have launched in the United States and ROW.

For a discussion of ongoing litigation related to XGEVA, see Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements.

TEPEZZA

Total TEPEZZA sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
TEPEZZA — U.S.	\$ 1,758	(4)%	\$ 1,835	*	\$ 441
TEPEZZA — ROW	145	*	16	*	7
Total TEPEZZA	<u>\$ 1,903</u>	<u>3 %</u>	<u>\$ 1,851</u>	<u>*</u>	<u>\$ 448</u>

* Change in excess of 100%

The increase in global TEPEZZA sales for 2025 was primarily driven by higher net selling price.

TEPEZZA was acquired on October 6, 2023 from our Horizon acquisition and generated \$1.9 billion and \$448 million in product sales for 2024 and 2023, respectively. As TEPEZZA was acquired on October 6, 2023, there were no recorded product sales in the period prior to the acquisition date.

BLINCYTO

Total BLINCYTO sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
BLINCYTO — U.S.	\$ 1,049	31 %	\$ 800	41 %	\$ 566
BLINCYTO — ROW	510	23 %	416	41 %	295
Total BLINCYTO	<u>\$ 1,559</u>	<u>28 %</u>	<u>\$ 1,216</u>	<u>41 %</u>	<u>\$ 861</u>

The increases in global BLINCYTO sales for 2025 and 2024 were driven by volume growth.

Nplate

Total Nplate sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Nplate — U.S.	\$ 1,027	6 %	\$ 970	(3)%	\$ 996
Nplate — ROW	497	2 %	486	1 %	481
Total Nplate	<u>\$ 1,524</u>	<u>5 %</u>	<u>\$ 1,456</u>	<u>(1)%</u>	<u>\$ 1,477</u>

Global Nplate sales for 2025 increased 5% and included U.S. government orders of \$90 million and \$128 million for 2025 and 2024, respectively. Excluding the U.S. government orders from this comparison, global Nplate sales increased 8% for 2025, driven by volume growth.

Global Nplate sales for 2024 decreased 1% and included U.S. government orders of \$128 million and \$286 million for 2024 and 2023, respectively. Excluding the U.S. government orders from this comparison, global Nplate sales increased 12% for 2024, driven by volume growth of 8% and higher net selling price of 6%.

TEZSPIRE

Total TEZSPIRE sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
TEZSPIRE — U.S.	\$ 1,478	52 %	\$ 972	71 %	\$ 567

The increases in TEZSPIRE sales for 2025 and 2024 were driven by volume growth. For 2026, we expect product sales for TEZSPIRE to follow the historical pattern of lower sales in the first quarter relative to subsequent quarters, as discussed above.

KYPROLIS

Total KYPROLIS sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
KYPROLIS — U.S.	\$ 913	(4)%	\$ 948	3 %	\$ 921
KYPROLIS — ROW	499	(10)%	555	15 %	482
Total KYPROLIS	<u>\$ 1,412</u>	<u>(6)%</u>	<u>\$ 1,503</u>	<u>7 %</u>	<u>\$ 1,403</u>

The decrease in global KYPROLIS sales for 2025 was primarily driven by lower volume due to increased competition.

The increase in global KYPROLIS sales for 2024 was driven by volume growth outside the United States.

Aranesp

Total Aranesp sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Aranesp — U.S.	\$ 416	8 %	\$ 386	(15)%	\$ 452
Aranesp — ROW	973	2 %	956	5 %	910
Total Aranesp	<u>\$ 1,389</u>	<u>4 %</u>	<u>\$ 1,342</u>	<u>(1)%</u>	<u>\$ 1,362</u>

The increase in global Aranesp sales for 2025 was driven by volume growth.

Global Aranesp sales for 2024 remained relatively unchanged as unfavorable changes to both estimated sales deductions and foreign currency exchange rates were offset by volume growth outside the United States.

KRYSTEXXA

Total KRYSTEXXA sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
KRYSTEXXA — U.S.	\$ 1,340	13 %	\$ 1,185	*	\$ 272

* Change in excess of 100%

The increase in KRYSTEXXA sales for 2025 was driven by volume growth and higher net selling price. For 2026, we expect product sales for KRYSTEXXA to follow the historical pattern of lower sales in the first quarter relative to subsequent quarters, as discussed above.

KRYSTEXXA was acquired on October 6, 2023 from our Horizon acquisition and generated \$1.2 billion and \$272 million in product sales for 2024 and 2023, respectively. As KRYSTEXXA was acquired on October 6, 2023, there were no recorded product sales in the period prior to the acquisition date.

Vectibix

Total Vectibix sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Vectibix — U.S.	\$ 604	16 %	\$ 519	13 %	\$ 461
Vectibix — ROW	571	9 %	526	1 %	523
Total Vectibix	<u>\$ 1,175</u>	<u>12 %</u>	<u>\$ 1,045</u>	<u>6 %</u>	<u>\$ 984</u>

The increase in global Vectibix sales for 2025 was primarily driven by volume growth.

The increase in global Vectibix sales for 2024 was driven by higher net selling price of 8% and volume growth of 4%, partially offset by unfavorable changes to foreign currency exchange rates.

Other products

Other product sales by geographic region were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
MVASI — U.S.	\$ 573	28 %	\$ 449	(12)%	\$ 511
MVASI — ROW	198	(29)%	278	(4)%	289
PAVBLU — U.S.	691	*	31	N/A	—
PAVBLU — ROW	9	N/A	—	N/A	—
UPLIZNA — U.S. ⁽¹⁾	528	68 %	314	*	60
UPLIZNA — ROW ⁽¹⁾	127	95 %	65	*	5
IMDELLTRA — U.S.	513	*	115	N/A	—
IMDYLLTRA — ROW	114	N/A	—	N/A	—
AMJEVITA — U.S.	48	(76)%	202	60 %	126
AMGEVITA — ROW	549	(2)%	559	12 %	500
TAVNEOS — U.S.	423	65 %	256	*	126
TAVNEOS — ROW	36	33 %	27	*	8
Neulasta — U.S.	359	13 %	318	(55)%	710
Neulasta — ROW	76	(33)%	113	(18)%	138
LUMAKRAS — U.S.	211	(1)%	214	9 %	197
LUMYKRAS — ROW	152	12 %	136	64 %	83
RAVICTI — U.S. ⁽¹⁾	337	(15)%	396	*	86
RAVICTI — ROW ⁽¹⁾	21	31 %	16	*	1
Parsabiv — U.S.	192	(5)%	203	(11)%	228
Parsabiv — ROW	161	5 %	153	14 %	134
Aimovig — U.S.	311	1 %	308	2 %	303
Aimovig — ROW	23	10 %	21	5 %	20
WEZLANA — U.S.	123	N/A	—	N/A	—
WEZENLA — ROW	150	*	27	N/A	—
PROCYSBI — U.S. ⁽¹⁾	233	5 %	221	*	49
PROCYSBI — ROW ⁽¹⁾	7	(13)%	8	*	1
Other — U.S. ⁽²⁾	895	(11)%	1,010	11 %	911
Other — ROW ⁽²⁾	203	7 %	190	(10)%	210
Total other product sales	<u>\$ 7,263</u>	<u>29 %</u>	<u>\$ 5,630</u>	<u>20 %</u>	<u>\$ 4,696</u>
Total U.S. — other products	\$ 5,437	35 %	\$ 4,037	22 %	\$ 3,307
Total ROW — other products	1,826	15 %	1,593	15 %	1,389
Total other product sales	<u>\$ 7,263</u>	<u>29 %</u>	<u>\$ 5,630</u>	<u>20 %</u>	<u>\$ 4,696</u>

* Change in excess of 100%

N/A = not applicable

⁽¹⁾ UPLIZNA, RAVICTI and PROCYSBI were acquired from our Horizon acquisition on October 6, 2023, and include product sales in the periods after the acquisition date.

⁽²⁾ Consists of product sales from (i) AVSOLA, KANJINTI, EPOGEN, RIABNI, BKEMV/BEKEMV, IMLYGIC, NEUPOGEN, Corlanor and Sensipar/Mimpara; and (ii) ACTIMMUNE, BUPHENYL, RAYOS, QUINSAIR, DUEXIS, VIMOVO and PENNSAID in the periods after our Horizon acquisition on October 6, 2023.

Operating expenses

Operating expenses were as follows (dollar amounts in millions):

	Year ended December 31, 2025	Change	Year ended December 31, 2024	Change	Year ended December 31, 2023
Cost of sales	\$ 12,037	(6)%	\$ 12,858	52 %	\$ 8,451
% of product sales	34.2 %		40.1 %		31.4 %
% of total revenues	32.8 %		38.5 %		30.0 %
Research and development	\$ 7,272	22 %	\$ 5,964	25 %	\$ 4,784
% of product sales	20.7 %		18.6 %		17.8 %
% of total revenues	19.8 %		17.8 %		17.0 %
Selling, general and administrative	\$ 7,050	(1)%	\$ 7,096	15 %	\$ 6,179
% of product sales	20.1 %		22.2 %		23.0 %
% of total revenues	19.2 %		21.2 %		21.9 %
Other	\$ 1,312	*	\$ 248	(72)%	\$ 879
Total operating expenses	\$ 27,671	6 %	\$ 26,166	29 %	\$ 20,293

* Change in excess of 100%

Cost of sales

Cost of sales decreased to 32.8% of total revenues for 2025, driven by lower amortization expense from acquisition-related assets, including the fair value step-up of inventory acquired from Horizon, and lower manufacturing costs, partially offset by higher profit share expense and changes in our sales mix. See Part IV—Note 4, Acquisition, to the Consolidated Financial Statements.

Cost of sales increased to 38.5% of total revenues for 2024, driven by higher amortization expense from Horizon acquisition-related assets and, to a lesser extent, higher profit share and royalty expense, partially offset by the prior year impact of the 2022 Puerto Rico tax law change that replaced an excise tax with an income tax beginning in 2023. For 2024, the unfavorable impact from product sales mix of certain Amgen products was offset by the favorable impact on product sales mix of the addition of acquired Horizon products.

Research and development

The Company groups all of its R&D activities and related expenditures into three categories: (i) Research and Early Pipeline, (ii) Later-Stage Clinical Programs and (iii) Marketed Product Support. These categories are described below:

Category	Description
Research and Early Pipeline	R&D expenses incurred in activities substantially in support of early research through the completion of Phase 1 clinical trials, including drug discovery, toxicology, pharmacokinetics and drug metabolism and process development
Later-Stage Clinical Programs	R&D expenses incurred in or related to Phase 2 and Phase 3 clinical programs intended to result in registration of a new product or a new indication for an existing product primarily in the United States or the EU
Marketed Product Support	R&D expenses incurred in support of the Company's marketed products that are authorized to be sold primarily in the United States or the EU. Includes clinical trials designed to gather information on product safety (certain of which may be required by regulatory authorities) and their product characteristics after regulatory approval has been obtained, as well as the costs of obtaining regulatory approval of a product in a new market after approval in either the United States or the EU has been obtained

R&D expense by category was as follows (in millions):

	Years ended December 31,		
	2025	2024	2023
Research and Early Pipeline	\$ 1,732	\$ 1,534	\$ 1,584
Later-Stage Clinical Programs	4,281	2,830	1,898
Marketed Product Support	1,259	1,600	1,302
Total R&D expense	\$ 7,272	\$ 5,964	\$ 4,784

The increase in R&D expense for 2025 was driven by investments in Later-Stage Clinical Programs, including those related to MariTide, and in Research and Early Pipeline, partially offset by lower spend in Marketed Product Support. This increase includes the impact of business development activities in 2025.

The increase in R&D expense for 2024 was driven by investments in Later-Stage Clinical Programs and Marketed Product Support, including Horizon-acquired programs.

We expect to continue to grow our spend on Later-Stage Clinical Programs as we advance our pipeline.

Selling, general and administrative

The decrease in SG&A expense for 2025 was driven by lower Horizon acquisition-related expenses and lower amortization expense from acquisition-related assets, partially offset by higher general and administrative expenses.

The increase in SG&A expense for 2024 was primarily driven by expenses from the acquired Horizon business and other commercial expenses, partially offset by lower acquisition-related expenses related to the Horizon acquisition incurred in 2024.

Other

Other operating expenses for 2025 included Otezla intangible asset impairment charges of \$1.2 billion. See Part IV—Note 13, Goodwill and other intangible assets, to the Consolidated Financial Statements.

Other operating expenses for 2024 included impairment charges associated with IPR&D intangible assets related to our Teneobio acquisition in 2021 and expenses related to cost-savings initiatives incurred in 2024.

Other operating expenses for 2023 included a net IPR&D intangible asset impairment charge for AMG 340 and expenses related to our restructuring plan that were both initiated and substantially completed in 2023.

Nonoperating expenses/income and income taxes

Nonoperating expenses/income and income taxes were as follows (dollar amounts in millions):

	Years ended December 31,		
	2025	2024	2023
Interest expense, net	\$ (2,755)	\$ (3,155)	\$ (2,875)
Other income, net	\$ 2,651	\$ 506	\$ 2,833
Provision for income taxes	\$ 1,265	\$ 519	\$ 1,138
Effective tax rate	14.1 %	11.3 %	14.5 %

Interest expense, net

The decrease in Interest expense, net, for 2025 was primarily due to lower average debt outstanding driven by deleveraging and, to a lesser extent, lower weighted-average fixed and floating interest rates on the debt. See Part IV—Note 16, Financing arrangements, to the Consolidated Financial Statements.

The increase in Interest expense, net, for 2024 was primarily due to higher average debt outstanding and higher weighted-average fixed and floating interest rates on the debt.

Other income, net

The increase in Other income, net, for 2025 was primarily due to current year net unrealized gains on equity investments, primarily BeOne, compared to net unrealized losses on equity investments in the prior year. See Part IV—Note 10, Investments, to the Consolidated Financial Statements.

The decrease in Other income, net, for 2024 was primarily due to net unrealized losses on equity investments in 2024 compared to net unrealized gains on equity investments in 2023, as well as reduced interest income as a result of lower average cash balances. The 2023 net unrealized gains on equity investments were principally composed of amounts recognized on our BeOne investment in the first quarter of 2023 as a result of a change from the equity method of accounting to recording this investment at fair value with changes in fair value recognized in earnings.

Income taxes

The increase in our effective tax rate for 2025 compared with 2024 was primarily due to a change in earnings mix, including the net unrealized gains on equity investments compared to net unrealized losses on equity investments in the prior year, partially offset by the prior-year deferred tax adjustments associated with U.S. tax on the earnings of our foreign subsidiaries and the current year Otezla intangible asset impairment charges and related tax impacts. See Part IV—Note 10, Investments, and Note 13, Goodwill and other intangible assets, to the Consolidated Financial Statements for additional information on our equity investments and Otezla impairment charges, respectively.

In 2021, the OECD reached an initial agreement to align countries on a minimum corporate tax rate and an expansion of the taxing rights of market countries. Select individual countries, including the United Kingdom, EU member countries and Singapore, have enacted the global minimum tax agreement that took effect starting in 2024. Singapore's enactment of the agreement, effective 2025, applies irrespective of the Company's incentive grant. On January 5, 2026, the OECD issued additional administrative guidance related to the global minimum tax agreement that exempts U.S. companies from extra territorial minimum taxes effective January 1, 2026. The new guidance did not impact our 2025 results. We are monitoring the potential 2026 impact of the guidance as jurisdictions may enact the new rules. See Part I, Item 1A. Risk Factors—*We could be subject to additional tax liabilities, including from an adverse outcome in our ongoing tax dispute with the IRS and other tax examinations, enactment of the OECD minimum corporate tax rate agreement and the adoption and interpretation of new tax legislation, including OB3. Such tax liabilities could adversely affect our profitability and results of operations.*

On July 4, 2025, OB3 was enacted in the United States. OB3 has various provisions, including the permanent extension of certain expiring provisions of the 2017 Tax Act, and modifications to the international tax framework. The legislation has multiple effective dates, with certain provisions effective in 2026 and beyond.

In 2017, we received an RAR and a modified RAR from the IRS for the years 2010–2012, proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico. We disagreed with the proposed adjustments and calculations, and in 2021, we filed a petition in the U.S. Tax Court to contest two duplicate Statutory Notices of Deficiency (Notices) for the years 2010–2012. The Notices seek to increase our U.S. taxable income for the years 2010–2012 by an amount that would result in additional federal tax of approximately \$3.6 billion plus interest. Any additional tax that could be imposed for the years 2010–2012 would be reduced by up to approximately \$900 million of repatriation tax previously accrued and paid on our foreign earnings.

In 2020, we received an RAR and a modified RAR from the IRS for the years 2013–2015, also proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico similar to those proposed for the years 2010–2012. We disagreed with the proposed adjustments and calculations, and in 2022, filed a petition in the U.S. Tax Court to contest a Notice for the years 2013–2015. The Notice seeks to increase our U.S. taxable income for the years 2013–2015 by an amount that would result in additional federal tax of approximately \$5.1 billion, plus interest, and asserts penalties of approximately \$2.0 billion. Any additional tax that could be imposed for the years 2013–2015 would be reduced by up to approximately \$2.2 billion of repatriation tax previously accrued and paid on our foreign earnings.

We firmly believe that the IRS positions set forth in the 2010–2012 and 2013–2015 Notices are without merit. We continue to contest the 2010–2012 and 2013–2015 Notices through the judicial process. The two cases were consolidated in the U.S. Tax Court in 2022. The trial began on November 4, 2024 and concluded on January 17, 2025. The parties filed opening post-trial briefs on June 13, 2025, and the Court held oral argument on July 16, 2025. The parties filed post-trial reply briefs on October 10, 2025. The Company expects a decision from the U.S. Tax Court no earlier than the second half of 2026.

We are currently under examination by the IRS for the years 2016–2018 with respect to issues similar to those for the 2010 through 2015 period. We expect that the IRS will begin its audit of 2019–2022 in the first half of 2026, and we believe

that it may seek to continue to audit similar issues related to the allocation of income between the United States and our foreign jurisdictions. In addition, we are under examination by a number of state and foreign tax jurisdictions.

Final resolution of these complex matters is not likely within the next 12 months. We continue to believe our accrual for income tax liabilities is appropriate based on past experience, interpretations of tax law, application of the tax law to our facts and judgments about potential actions by tax authorities; however, due to the complexity of the provision for income taxes and uncertain resolution of these matters, the ultimate outcome of any tax matters may result in payments substantially greater than amounts accrued and could have a material adverse impact on our consolidated financial statements.

See Part I, Item 1A. Risk Factors—*We could be subject to additional tax liabilities, including from an adverse outcome in our ongoing tax dispute with the IRS and other tax examinations, enactment of the OECD minimum corporate tax rate agreement and the adoption and interpretation of new tax legislation, including OB3. Such tax liabilities could adversely affect our profitability and results of operations*; Part II, Item 7. Management’s Discussion and Analysis of Financial Condition and Results of Operations—Critical Accounting Policies and Estimates—*Income taxes*; and Part IV—Note 7, Income taxes, to the Consolidated Financial Statements for further discussion.

Financial condition, liquidity and capital resources

Selected financial data was as follows (in millions):

	December 31,	
	2025	2024
Cash and cash equivalents	\$ 9,129	\$ 11,973
Total assets	\$ 90,586	\$ 91,839
Current portion of long-term debt	\$ 4,599	\$ 3,550
Long-term debt	\$ 50,005	\$ 56,549
Stockholders’ equity	\$ 8,658	\$ 5,877

Cash and cash equivalents

Our balance of cash and cash equivalents was \$9.1 billion as of December 31, 2025. The primary objective of our investment portfolio is to maintain safety of principal, prudent levels of liquidity and acceptable levels of risk. Our investment policy limits interest-bearing security investments to certain types of debt and money market instruments issued by institutions with primarily investment-grade credit ratings, and it places restrictions on maturities and concentration by asset class and issuer.

Capital allocation

Consistent with the objective to optimize our capital structure, we deploy our accumulated cash balances in a strategic manner and consider a number of alternatives, including investments in innovation both internally and externally (including investments that expand our portfolio of products in areas of therapeutic interest), capital expenditures, repayment of debt, payment of dividends and stock repurchases.

We intend to continue investing in our business while returning capital to stockholders through the payment of cash dividends and stock repurchases. This reflects our desire to optimize our cost of capital and our confidence in the future cash flows of our business. The timing and amount of future dividends and stock repurchases will vary based on a number of factors, including future capital requirements for strategic transactions, debt levels and debt service requirements, our credit rating, availability of financing on acceptable terms, changes to applicable tax laws or corporate laws, changes to our business model and periodic determination by our Board of Directors that cash dividends and/or stock repurchases are in the best interests of stockholders and are in compliance with applicable laws and the Company’s agreements. In addition, the timing and amount of stock repurchases may also be affected by our overall level of cash, stock price and blackout periods, during which we are restricted from repurchasing stock. The manner of stock repurchases may include block purchases, tender offers, accelerated share repurchases and market transactions.

The Board of Directors declared quarterly cash dividends of \$2.38, \$2.25 and \$2.13 per share of common stock paid in 2025, 2024 and 2023, respectively, reflecting year-over-year increases of 6% for both 2025 and 2024. In December 2025, the Board of Directors declared a cash dividend of \$2.52 per share of common stock for the first quarter of 2026, an increase of 6% over the same period in the prior year, which will be paid in March 2026.

We also return capital to stockholders through our stock repurchase program. During 2025 and 2023, we did not repurchase shares under the stock repurchase program. During 2024, we repurchased \$200 million of common stock under the stock repurchase program. As of December 31, 2025, \$6.8 billion of authorization remained available under the stock repurchase program.

As a result of stock repurchases and quarterly dividend payments, we had an accumulated deficit as of December 31, 2025 and 2024. Our accumulated deficit is not anticipated to affect our future ability to operate, repurchase stock, pay dividends or repay our debt given our expected continued profitability and strong financial position.

We believe that existing funds, cash generated from operations and existing sources of and access to financing are adequate to satisfy our needs for working capital, capital expenditure and debt service requirements, pay dividends and repurchase stock, and other business initiatives we plan to strategically pursue, including acquisitions and licensing activities. We anticipate that our liquidity needs can be met through a variety of sources, including cash provided by operating activities, borrowings through commercial paper and/or syndicated credit facilities, and access to other domestic and foreign debt markets and equity markets. See Part I, Item 1A. Risk Factors—*Global economic conditions may negatively affect us and may magnify certain risks that affect our business.*

Financing arrangements

To help meet our liquidity requirements, we have entered into various financing arrangements. The noncurrent portions of our long-term borrowings as of December 31, 2025 and 2024, were \$50.0 billion and \$56.5 billion, respectively. The carrying values of our long-term borrowings are net of fair value adjustments for interest rate swaps and unamortized discounts, premiums and offering costs. As of December 31, 2025, S&P, Moody's and Fitch assigned credit ratings to our outstanding senior notes of BBB+, Baa1 and BBB+, respectively, which are considered investment grade. Unfavorable changes to these ratings may have an adverse impact on future financings.

During 2023, in connection with our acquisition of Horizon, we issued \$24.0 billion of debt composed of eight series of notes and borrowed \$4.0 billion under a term loan credit agreement, of which \$1.8 billion of borrowings was outstanding as of December 31, 2025.

During 2025, we retired \$6.0 billion of debt, consisting of \$5.0 billion of debt repayments and \$1.0 billion of debt repurchases. The debt repurchases were completed for an aggregate cost of \$683 million and resulted in a \$264 million gain on extinguishment of debt. We periodically consider the repurchase of our debt when conditions are favorable. Gains on extinguishment of debt are recorded in Other income, net in the Consolidated Statements of Income.

During 2024, we retired \$4.5 billion of debt, consisting of \$3.6 billion of debt repayments, of which \$2.2 billion related to repayments on our term loans, and \$875 million of debt repurchases. The debt repurchases were completed for an aggregate cost of \$659 million and resulted in a \$215 million gain on extinguishment of debt.

During 2023, we retired \$2.3 billion of debt, consisting of \$1.5 billion of debt repayments and \$881 million of debt repurchases. The debt repurchases were completed for an aggregate cost of \$647 million and resulted in a \$225 million gain on extinguishment of debt.

To achieve a desired mix of fixed-rate and floating-rate debt, we entered into interest rate swap contracts that effectively converted a fixed-rate interest coupon for certain of our debt issuances to a floating, SOFR-based coupon over the terms of the respective notes. These interest rate swap contracts qualify and are designated as fair value hedges. As of both December 31, 2025 and 2024, we had interest rate swap contracts with an aggregate notional amount of \$6.7 billion.

To hedge our exposure to foreign currency exchange rate risk associated with certain of our long-term notes denominated in foreign currencies, we entered into cross-currency swap contracts, which effectively converted the interest payments and principal repayment of the respective notes from euros and pounds sterling to U.S. dollars. These cross-currency swap contracts qualify and are designated as cash flow hedges. As of both December 31, 2025 and 2024, we had cross-currency swap contracts with an aggregate notional amount of \$2.7 billion.

In 2025, we increased the capacity of our commercial paper program from \$2.5 billion to \$4.0 billion, which allows us to issue up to \$4.0 billion of unsecured commercial paper to fund working capital needs. We did not issue any commercial paper during 2025, 2024 or 2023, and no commercial paper was outstanding as of December 31, 2025 and 2024.

In 2023, we amended and restated our syndicated, unsecured, revolving credit agreement, under which we may borrow up to \$4.0 billion for general corporate purposes, including as a liquidity backstop for our commercial paper program. The commitments under the revolving credit agreement may be increased by up to \$1.25 billion with the agreement of the banks. Each bank that is a party to the agreement has an initial commitment term of five years. This term may be extended for up to two additional one-year periods with the agreement of the banks. Annual commitment fees for this agreement are 0.09% of the

unused portion of the facility based on our current credit rating. Generally, we would be charged interest for any amounts borrowed under this facility, based on our current credit rating, at (i) SOFR plus 1.01% or (ii) the highest of (A) the administrative agent bank base commercial lending rate, (B) the overnight federal funds rate plus 0.50% or (C) one-month SOFR plus 1.1%. As of December 31, 2025 and 2024, no amounts were outstanding under this facility.

Also in 2023, we filed a shelf registration statement with the SEC that allows us to issue unspecified amounts of debt securities; common stock; preferred stock; warrants to purchase debt securities, common stock, preferred stock or depositary shares; rights to purchase common stock or preferred stock; securities purchase contracts; securities purchase units; and depositary shares. Under this shelf registration statement, all of the securities available for issuance may be offered from time to time, with terms to be determined at the time of issuance. This shelf registration statement expired in February 2026, and our Board has approved a new shelf registration statement to replace it.

Certain of our financing arrangements contain nonfinancial covenants. In addition, our revolving credit agreement and term loan credit agreement include a financial covenant that requires us to maintain a specified minimum interest coverage ratio of (i) the sum of consolidated net income, interest expense, provision for income taxes, depreciation expense, amortization expense, unusual or nonrecurring charges and other noncash items (consolidated earnings before interest, taxes, depreciation and amortization) to (ii) Consolidated Interest Expense, each as defined and described in the respective agreements. We were in compliance with all applicable covenants under these arrangements as of December 31, 2025.

These financing arrangements are more fully discussed in Part IV—Note 16, Financing arrangements, and Note 19, Derivative instruments, to the Consolidated Financial Statements.

Cash flows

Our summarized cash flow activity was as follows (in millions):

	Years ended December 31,		
	2025	2024	2023
Net cash provided by operating activities	\$ 9,958	\$ 11,490	\$ 8,471
Net cash used in investing activities	\$ (1,943)	\$ (1,046)	\$ (26,204)
Net cash (used in) provided by financing activities	\$ (10,859)	\$ (9,415)	\$ 21,048

Operating

Cash provided by operating activities has been and is expected to continue to be our primary recurring source of funds. Cash provided by operating activities decreased in 2025 as compared to 2024 due to the timing of working capital items primarily driven by higher collections in the fourth quarter of 2024, partially offset by higher net income after adjustments for noncash items and lower interest payments.

Cash provided by operating activities increased in 2024 as compared to 2023 due to higher net income after adjustments for noncash items and the timing of working capital items primarily driven by higher collections in the fourth quarter of 2024.

Investing

Cash used in investing activities during 2025 and 2024 was primarily due to \$1.9 billion and \$1.1 billion, respectively, of capital expenditures, including construction costs for new plants and expansion of manufacturing capacity.

Cash used in investing activities during 2023 was primarily due to \$27.0 billion of net cash used for the purchase of Horizon and \$1.1 billion of capital expenditures, partially offset by net cash inflows related to marketable securities of \$1.7 billion.

We currently estimate 2026 investments in capital projects to be approximately \$2.6 billion. A majority of the increase in expenditures relates to construction costs for new plants and expansion of manufacturing capacity to enable supply of products and product candidates.

Financing

Cash used in financing activities during 2025 was primarily due to the repayment and extinguishment of debt of \$5.0 billion and \$683 million, respectively, and the payment of dividends of \$5.1 billion.

Cash used in financing activities during 2024 was primarily due to the payment of dividends of \$4.8 billion, the repayment and extinguishment of debt of \$3.6 billion and \$659 million, respectively, and payments to repurchase common stock of \$200 million.

Cash provided by financing activities during 2023 was primarily due to net proceeds from long-term debt issuances of \$27.8 billion primarily in connection with the acquisition of Horizon, partially offset by the payment of dividends of \$4.6 billion and the repayment and extinguishment of debt of \$1.5 billion and \$647 million, respectively.

See Part IV—Note 10, Investments; Note 16, Financing arrangements; and Note 17, Stockholders' equity, to the Consolidated Financial Statements.

Capital requirements

We have material cash requirements to pay third parties under various contractual obligations discussed below.

We are obligated to pay interest and repay principal under our various financing arrangements, including amounts under interest rate swap and cross-currency swap contracts related to certain of our long-term debt obligations. For information on scheduled debt maturities and payments under derivative contracts associated with our long-term debt obligations, see Part IV—Note 16, Financing arrangements, and Note 19, Derivative instruments, to the Consolidated Financial Statements.

We are obligated to make payments for operating leases, including rental commitments on unoccupied leases and leases that have not yet commenced. For information on these obligations, see Part IV—Note 14, Leases, to the Consolidated Financial Statements.

As of December 31, 2025, we have purchase obligations of approximately \$6.9 billion primarily related to (i) R&D commitments (including those related to clinical trials) for new and existing products, (ii) capital expenditures and (iii) open purchase orders for the acquisition of goods and services in the ordinary course of business. Most of these obligations are expected to be paid within one year, and payment of certain of these amounts may be reduced based on certain future events.

In addition to the purchase obligations noted above and upon the achievement of various development, regulatory and commercial milestones for agreements we have entered into with third parties, we are contractually obligated to pay additional amounts that, in the aggregate, are significant. These payments are contingent upon the occurrence of various future events, substantially all of which have a high degree of uncertainty of occurring, and any resulting cash requirements are managed through our operational budgeting processes. Except with respect to the fair value of the contingent consideration of approximately \$161 million as of December 31, 2025, these obligations are not recorded on our Consolidated Balance Sheets. As of December 31, 2025, the maximum amount that may be payable in the future for agreements we have entered into with third parties is approximately \$7.2 billion.

We have recorded liabilities for UTBs that, because of their nature, have a high degree of uncertainty regarding the timing of future cash payment and other events that extinguish these liabilities. See Part IV—Note 7, Income taxes, to the Consolidated Financial Statements.

Critical accounting policies and estimates

The preparation of our consolidated financial statements in conformity with GAAP requires management to make estimates and assumptions that affect the amounts reported in the financial statements and the notes to the financial statements. Some of those judgments can be subjective and complex, and therefore, actual results could differ materially from those estimates under different assumptions or conditions. Our significant accounting policies are included in Part IV—Note 1, Summary of significant accounting policies, to the Consolidated Financial Statements. The following are considered critical to our consolidated financial statements because they require the most difficult, subjective or complex judgments, often because of the need to make estimates about matters that are inherently uncertain.

Product sales and sales deductions

Revenue from product sales is recognized upon transfer of control of a product to a customer, generally upon delivery, based on an amount that reflects the consideration to which we expect to be entitled, net of accruals for estimated rebates, wholesaler chargebacks, discounts and other deductions (collectively, sales deductions) established at the time of sale.

We analyze the adequacy of our accruals for sales deductions quarterly. Amounts accrued for sales deductions are adjusted when trends or significant events indicate that adjustment is appropriate. Accruals are also adjusted to reflect actual results. Amounts recorded in Accrued liabilities in the Consolidated Balance Sheets for sales deductions were as follows (in millions):

	Rebates	Chargebacks	Other deductions	Total
Balance as of December 31, 2022	\$ 4,879	\$ 849	\$ 258	\$ 5,986
Additions ⁽¹⁾	263	24	39	326
Amounts charged against product sales	14,328	13,349	2,533	30,210
Payments	(13,634)	(13,125)	(2,492)	(29,251)
Balance as of December 31, 2023	5,836	1,097	338	7,271
Amounts charged against product sales	17,404	14,882	3,060	35,346
Payments	(16,423)	(14,817)	(2,972)	(34,212)
Balance as of December 31, 2024	6,817	1,162	426	8,405
Amounts charged against product sales	21,697	16,988	3,298	41,983
Payments	(19,675)	(16,852)	(3,255)	(39,782)
Balance as of December 31, 2025	\$ 8,839	\$ 1,298	\$ 469	\$ 10,606

⁽¹⁾ Represents sales deductions assumed from the Horizon acquisition.

For the years ended December 31, 2025, 2024 and 2023, total sales deductions were 54%, 52% and 53% of gross product sales, respectively. The increase in the total sales deductions balance as of December 31, 2025, compared with December 31, 2024, was primarily driven by an increase in gross sales and timing of payments. Included in the amounts are immaterial net adjustments related to prior-year sales due to changes in estimates.

In the United States, we use wholesalers as the principal means of distributing our products to healthcare providers such as physicians or their clinics, dialysis centers, hospitals and pharmacies. Products we sell in Europe are distributed principally to hospitals and/or wholesalers depending on the distribution practice in each country where the products are sold. We monitor the inventory levels of our products at our wholesalers by using data from our wholesalers and other third parties, and we believe wholesaler inventories have been maintained at appropriate levels (generally two to three weeks) given end-user demand. Accordingly, historical fluctuations in wholesaler inventory levels have not significantly affected our method of estimating sales deductions.

Accruals for sales deductions are based primarily on estimates of the amounts earned or to be claimed on the related sales. These estimates take into consideration current contractual and statutory requirements, specific known market events and trends, internal and external historical data and forecasted customer buying patterns. Sales deductions are substantially product specific and therefore, for any given year, can be affected by the mix of products sold.

Rebates include primarily amounts paid to payers and providers in the United States, including those paid to state Medicaid programs and those related to the IRA, and are based on contractual arrangements or statutory requirements that vary by product, by payer and by individual payer plans. As we sell products, we estimate the amount of rebate we will pay based on

the product sold, contractual terms, estimated patient population, historical experience and wholesaler inventory levels; and we accrue these rebates in the period the related sales are recorded. We then adjust the rebate accruals as more information becomes available and to reflect actual claims experience. Estimating such rebates is complicated, in part because of the time delay between the date of sale and the actual settlement of the liability. We believe the methodology we use to accrue for rebates is reasonable and appropriate given current facts and circumstances, but actual results may differ.

Wholesaler chargebacks relate to our contractual agreements to sell products to healthcare providers in the United States at fixed prices that are lower than the prices we charge wholesalers. When healthcare providers purchase our products through wholesalers at these reduced prices, wholesalers charge us for the difference between their purchase prices and the contractual prices between Amgen and the healthcare providers. The provision for chargebacks is based on expected sales by our wholesaler customers to healthcare providers. Accruals for wholesaler chargebacks are less difficult to estimate than rebates are, and they closely approximate actual results because chargeback amounts are fixed at the date of purchase by the healthcare providers and because we generally settle the liability for these deductions within a few weeks.

Income taxes

We provide for income taxes based on pretax income and applicable tax rates in the various jurisdictions in which we operate.

We recognize the tax benefit from an uncertain tax position only if it is more likely than not that the tax position will be sustained upon examination by tax authorities based on the technical merits of the position. The tax benefit recognized in the consolidated financial statements for a particular tax position is based on the largest benefit that is more likely than not to be realized. The amount of UTBs is adjusted as appropriate for changes in facts and circumstances, such as significant amendments to existing tax law, new regulations or interpretations by tax authorities, new information obtained during a tax examination or resolution of an examination. We believe our estimates for uncertain tax positions are appropriate and sufficient for any assessments that may result from examinations of our tax returns. We recognize both accrued interest and penalties, when appropriate, related to UTBs in income tax expense. See Part IV—Note 7, Income taxes, to the Consolidated Financial Statements.

Certain items are included in our tax return at different times than they are reflected in the financial statements, and they cause temporary differences between the tax bases of assets and liabilities and their reported amounts. Such temporary differences create deferred tax assets and liabilities. Deferred tax assets are generally items that can be used as tax deductions or credits in tax returns in future years but for which we have already recorded the tax benefit in the consolidated financial statements. We establish valuation allowances against our deferred tax assets when the amount of expected future taxable income is not likely to support the use of the deduction or credit. Deferred tax liabilities are either (i) tax expenses recognized in the consolidated financial statements for which payment has been deferred, (ii) expenses for which we have already taken a deduction on the tax return but have not yet recognized in the consolidated financial statements or (iii) liabilities for the difference between the book basis and the tax basis of the intangible assets acquired in many business combinations, because future expenses associated with these assets most often will not be tax deductible.

Amgen is subject to current U.S. tax on the earnings of our foreign subsidiaries. We previously established deferred taxes related to this U.S. tax, which requires us to recognize deferred taxes for temporary basis differences expected to reverse and be subject to this tax in future years. These are ongoing adjustments that are likely to occur in future periods.

We are a vertically integrated enterprise with operations in the United States and various foreign jurisdictions. In the jurisdictions where we conduct operations, we are subject to income tax based on the tax laws and principles of such jurisdictions and on the functions, risks and activities performed therein. Our pretax income is therefore attributed to domestic or foreign sources based on the operations performed and the risks assumed in each location, as well as on the tax laws and principles of the respective taxing jurisdictions. For example, we conduct significant operations in Puerto Rico, a territory of the United States that is treated as a foreign jurisdiction for U.S. tax purposes, pertaining to manufacturing, distribution and other related functions to meet our worldwide product demand. Income from our operations in Puerto Rico is subject to tax incentive grants through 2050.

In 2017, we received an RAR and a modified RAR from the IRS for the years 2010–2012, proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico. We disagreed with the proposed adjustments and calculations, and in 2021, filed a petition in the U.S. Tax Court to contest two duplicate Statutory Notices of Deficiency (Notices) for the years 2010–2012. The Notices seek to increase our U.S. taxable income for the years 2010–2012 by an amount that would result in additional federal tax of approximately \$3.6 billion plus interest. Any additional tax that could be imposed for the years 2010–2012 would be reduced by up to approximately \$900 million of repatriation tax previously accrued and paid on our foreign earnings.

In 2020, we received an RAR and a modified RAR from the IRS for the years 2013–2015, also proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico similar to those proposed for the years 2010–2012. We disagreed with the proposed adjustments and calculations, and in 2022, filed a petition in the U.S. Tax Court to contest a Notice for the years 2013–2015. The Notice seeks to increase our U.S. taxable income for the years 2013–2015 by an amount that would result in additional federal tax of approximately \$5.1 billion, plus interest, and asserts penalties of approximately \$2.0 billion. Any additional tax that could be imposed for the years 2013–2015 would be reduced by up to approximately \$2.2 billion of repatriation tax previously accrued and paid on our foreign earnings.

We firmly believe that the IRS positions set forth in the 2010–2012 and 2013–2015 Notices are without merit. We continue to contest the 2010–2012 and 2013–2015 Notices through the judicial process. The two cases were consolidated in the U.S. Tax Court in 2022. The trial began on November 4, 2024, and concluded on January 17, 2025. The parties filed opening post-trial briefs on June 13, 2025, and the Court held oral argument on July 16, 2025. The parties filed post-trial reply briefs on October 10, 2025. The Company expects a decision from the U.S. Tax Court no earlier than the second half of 2026.

We are currently under examination by the IRS for the years 2016–2018 with respect to issues similar to those for the 2010 through 2015 period. We expect that the IRS will begin its audit of 2019–2022 in the first half of 2026, and we believe that it may seek to continue to audit similar issues related to the allocation of income between the United States and our foreign jurisdictions. In addition, we are under examination by a number of state and foreign tax jurisdictions.

Final resolution of these complex matters is not likely within the next 12 months. We continue to believe our accrual for income tax liabilities is appropriate based on past experience, interpretations of tax law, application of the tax law to our facts and judgments about potential actions by tax authorities; however, due to the complexity of the provision for income taxes and uncertain resolution of these matters, the ultimate outcome of any tax matters may result in payments substantially greater than amounts accrued and could have a material adverse impact on our consolidated financial statements.

See Part I, Item 1A. Risk Factors—*We could be subject to additional tax liabilities, including from an adverse outcome in our ongoing tax dispute with the IRS and other tax examinations, enactment of the OECD minimum corporate tax rate agreement and the adoption and interpretation of new tax legislation, including OB3. Such tax liabilities could adversely affect our profitability and results of operations*; Part II, Item 7. Management’s Discussion and Analysis of Financial Condition and Results of Operations—Results of Operations—Income taxes; and Part IV—Note 7, Income taxes, to the Consolidated Financial Statements for further discussion.

Our operations are subject to the tax laws, regulations and administrative practices of the United States, the U.S. territory of Puerto Rico, U.S. state jurisdictions and other countries in which we do business. Significant changes in these rules could have a material adverse effect on our results of operations. See Part I, Item 1A. Risk Factors—*We could be subject to additional tax liabilities, including from an adverse outcome in our ongoing tax dispute with the IRS and other tax examinations, enactment of the OECD minimum corporate tax rate agreement and the adoption and interpretation of new tax legislation, including OB3. Such tax liabilities could adversely affect our profitability and results of operations*.

Contingencies

In the ordinary course of business, we are involved in various legal proceedings, government investigations and other matters such as intellectual property disputes, contractual disputes and class action suits that are complex in nature and have outcomes that are difficult to predict. We describe our legal proceedings and other matters that are significant or that we believe could become significant in Part IV—Note 20, Contingencies and commitments, to the Consolidated Financial Statements. We record accruals for loss contingencies to the extent that we conclude it is probable that a liability has been incurred and the amount of the related loss can be reasonably estimated. We evaluate, on a quarterly basis, developments in legal proceedings and other matters that could cause an increase or decrease in the amount of the liability that has been accrued previously.

While it is not possible to accurately predict or determine the eventual outcomes of these items, an adverse determination in one or more of these items currently pending could have a material adverse effect on our consolidated results of operations, financial position or cash flows.

Valuation of assets and liabilities in connection with acquisitions

We have acquired and continue to acquire intangible assets in connection with business combinations and asset acquisitions. These intangible assets consist primarily of technology associated with currently marketed human therapeutic products and IPR&D product candidates. Discounted cash flow models are typically used to determine the fair values of these intangible assets for purposes of allocating consideration paid to the net assets acquired in an acquisition. See Part IV—Note 4, Acquisition, to the Consolidated Financial Statements. These models require the use of significant estimates and assumptions, including but not limited to:

- determining the timing and expected costs to complete in-process projects, taking into account the stage of completion at the acquisition date;
- projecting the probability and timing of obtaining marketing approval from the FDA and other regulatory agencies for product candidates;
- estimating the timing of and future net cash flows from product sales resulting from completed products and in-process projects; and
- developing appropriate discount rates to calculate the present values of the cash flows.

Significant estimates and assumptions are also required to determine the business combination date fair values of any contingent consideration obligations incurred in connection with business combinations. In addition, we must revalue these obligations each subsequent reporting period until the related contingencies are resolved and record changes in their fair values in earnings. The acquisition date fair values of contingent consideration obligations incurred or assumed in the acquisitions were determined using a combination of valuation techniques. Significant estimates and assumptions required for these valuations included but were not limited to the timing and probability of achieving regulatory milestones, product sales projections under various scenarios and discount rates used to calculate the present value of the required payments. These estimates and assumptions are required to be updated in order to revalue these contingent consideration obligations each reporting period. Accordingly, subsequent changes in underlying facts and circumstances could result in changes in these estimates and assumptions, which could have a material impact on the estimated future fair values of these obligations.

We believe the fair values used to record intangible assets acquired and contingent consideration obligations incurred in connection with business combinations and asset acquisitions are based on reasonable estimates and assumptions given the facts and circumstances as of the related valuation dates.

Impairment of long-lived assets

We review the carrying value of our finite-lived intangible assets for impairment whenever events or changes in circumstances indicate that the carrying amount of an asset may not be recoverable. If such circumstances exist, an estimate of undiscounted future cash flows to be generated by the long-lived asset is compared with the carrying value to determine whether an impairment exists. If an asset is determined to be impaired, the loss is measured based on the difference between the asset's fair value and its carrying value.

Indefinite-lived intangible assets, composed of IPR&D projects acquired in a business combination that have not reached technological feasibility or that lack regulatory approval at the time of acquisition, are reviewed for impairment annually, whenever events or changes in circumstances indicate that the carrying amount may not be recoverable and upon establishment of technological feasibility or regulatory approval. We test for impairment by comparing the fair value of the asset to its carrying value. If the asset's carrying value exceeds its fair value, an impairment charge is recorded for the difference, and its carrying value is reduced accordingly.

Estimating future cash flows of an IPR&D product candidate for purposes of an impairment analysis requires us to make significant estimates and assumptions regarding the amount and timing of costs to complete the project and the amount, timing and probability of achieving revenues from the completed product similar to how the acquisition date fair value of the project was determined, as described above. There are often major risks and uncertainties associated with IPR&D projects as we are required to obtain regulatory approvals in order to be able to market these products. Such approvals require completing clinical trials that demonstrate a product candidate is safe and effective. Consequently, the eventual realized value of the acquired IPR&D project may vary from its fair value at the date of acquisition, and IPR&D impairment charges may occur in future periods which could have a material adverse effect on our results of operations.

We believe our estimations of future cash flows used for assessing impairment of long-lived assets are based on reasonable assumptions given the facts and circumstances as of the related dates of the assessments.

Recently issued accounting standards

See Part IV—Note 1, Summary of significant accounting policies, to the Consolidated Financial Statements for a discussion of recently issued accounting pronouncements.

Item 7A. QUANTITATIVE AND QUALITATIVE DISCLOSURES ABOUT MARKET RISK

We are exposed to market risks that may result from changes in interest rates, foreign currency exchange rates and prices of equity instruments as well as changes in general economic conditions in the countries where we conduct business. To reduce certain of these risks, we enter into various types of foreign currency and interest rate derivative hedging transactions as part of our risk management program. We do not use derivatives for speculative trading purposes.

In the discussion that follows, we assumed a hypothetical change in interest rates of 100 basis points from those as of December 31, 2025 and 2024. Except as noted below, we also assumed a hypothetical 20% change in foreign currency exchange rates against the U.S. dollar based on its position relative to other currencies as of December 31, 2025 and 2024.

Interest-rate-sensitive financial instruments

Our portfolio of available-for-sale investments as of December 31, 2025 and 2024, was composed almost entirely of U.S. Treasury securities and money market mutual funds. The fair values of our available-for-sale investments were \$8.5 billion and \$11.5 billion as of December 31, 2025 and 2024, respectively. Duration is a sensitivity measure that can be used to approximate the change in the value of a security that will result from a 100 basis point change in interest rates. Applying a duration model, a hypothetical 100 basis point increase in interest rates as of December 31, 2025 and 2024, would not have resulted in a material reduction in the fair values of these securities. In addition, a hypothetical 100 basis point decrease in interest rates as of December 31, 2025 and 2024, would not result in a material effect on income in the respective ensuing year.

As of December 31, 2025, we had outstanding notes with an aggregate carrying value of \$52.8 billion and an aggregate fair value of \$51.0 billion. As of December 31, 2024, we had outstanding notes with an aggregate carrying value of \$58.3 billion and an aggregate fair value of \$54.9 billion. Our outstanding notes were composed of debt with fixed interest rates. Changes in interest rates do not affect interest expense on fixed-rate debt. Changes in interest rates would, however, affect the fair values of fixed-rate debt. A hypothetical 100 basis point decrease in interest rates relative to interest rates as of December 31, 2025 and 2024, would have resulted in increases of \$4.4 billion and \$4.7 billion, respectively, in the aggregate fair values of our outstanding debt on these dates. The sensitivity analysis of the notes does not consider the impact that hypothetical changes in interest rates would have on related interest rate swap contracts and cross-currency swap contracts, discussed below. In addition, the analysis above does not include our term loans, which had a carrying value of \$1.8 billion at both December 31, 2025 and 2024. The fair value of our term loan is approximated at its carrying value as this debt instrument bears interest at a floating rate.

To achieve a desired mix of fixed-rate and floating-rate debt, we entered into interest rate swap contracts that qualified and were designated for accounting purposes as fair value hedges for certain of our fixed-rate debt. These interest rate swap contracts effectively converted a fixed-rate interest coupon to a floating-rate SOFR-based coupon over the terms of the respective notes. Interest rate swap contracts with an aggregate notional amount of \$6.7 billion were outstanding as of both December 31, 2025 and 2024. A hypothetical 100 basis point increase in interest rates relative to interest rates as of December 31, 2025 and 2024, would have resulted in reductions in fair values of approximately \$240 million and \$220 million, respectively, on our interest rate swap contracts on these dates. Analysis of the interest rate swap contracts does not consider the impact that hypothetical changes in interest rates would have on the related fair values of debt that these interest-rate-sensitive instruments were designed to offset.

As of both December 31, 2025 and 2024, we had outstanding cross-currency swap contracts with an aggregate notional amount of \$2.7 billion that hedge our foreign-currency-denominated debt and related interest payments. These contracts effectively convert interest payments and principal repayment of this debt to U.S. dollars from euros and pounds sterling and are designated for accounting purposes as cash flow hedges. A hypothetical 100 basis point adverse movement in interest rates relative to interest rates as of December 31, 2025 and 2024, would not have a material effect on the fair values of our cross-currency swap contracts.

Foreign-currency-sensitive financial instruments

Our international operations are affected by fluctuations in the value of the U.S. dollar compared with foreign currencies, predominantly the euro. Increases and decreases in our international product sales from movements in foreign currency exchange rates are partially offset by corresponding increases or decreases in our international operating expenses. Increases and decreases in our foreign-currency-denominated assets from movements in foreign currency exchange rates are partially offset by corresponding increases or decreases in our foreign-currency-denominated liabilities. To further reduce our net exposure to foreign currency exchange rate fluctuations on our results of operations, we enter into foreign currency forward and cross-currency swap contracts.

As of December 31, 2025, we had outstanding euro- and pound-sterling-denominated debt with both a principal carrying value and a fair value of \$2.5 billion. As of December 31, 2024, we had outstanding euro- and pound-sterling-denominated debt with both a principal carrying value and a fair value of \$2.2 billion. A hypothetical 20% adverse movement in foreign currency exchange rates compared with the U.S. dollar relative to exchange rates as of December 31, 2025, would have resulted in an increase in fair value of this debt of approximately \$490 million on this date and a reduction in income in the ensuing year of approximately \$490 million. A hypothetical 20% adverse movement in foreign currency exchange rates compared with the U.S. dollar relative to exchange rates as of December 31, 2024, would have resulted in an increase in fair value of this debt of \$440 million on this date and a reduction in income in the ensuing year of \$450 million. The impact on income from these hypothetical changes in foreign currency exchange rates would be substantially offset by the impact such changes would have on related cross-currency swap contracts, which are in place for the related foreign-currency-denominated debt.

We have cross-currency swap contracts that are designated as cash flow hedges of our debt denominated in euros and pounds sterling with an aggregate notional amount of \$2.7 billion as of both December 31, 2025 and 2024. A hypothetical 20% adverse movement in foreign currency exchange rates compared with the U.S. dollar relative to exchange rates on these dates would have resulted in reductions in the fair values of these contracts of approximately \$500 million and \$450 million on these dates, respectively. The impact of this hypothetical adverse movement in foreign currency exchange rates on ensuing years' income from these contracts would be fully offset by corresponding hypothetical changes in the carrying amounts of the related hedged debt.

We enter into foreign currency forward contracts that are designated for accounting purposes as cash flow hedges of certain anticipated foreign currency transactions. As of December 31, 2025, the fair values of these contracts were a \$195 million asset and a \$213 million liability. As of December 31, 2024, the fair values of these contracts were a \$420 million asset and an \$8 million liability. As of December 31, 2025, we had primarily euro-based open foreign currency forward contracts with an aggregate notional amount of \$7.8 billion. As of December 31, 2024, we had primarily euro-based open foreign currency forward contracts with an aggregate notional amount of \$7.2 billion. With regard to foreign currency forward contracts that were open as of December 31, 2025, a hypothetical 20% adverse movement in foreign currency exchange rates compared with the U.S. dollar relative to exchange rates as of December 31, 2025, would have resulted in a reduction in fair value of these contracts of approximately \$1.5 billion on this date and in the ensuing year, a reduction in income of approximately \$800 million. With regard to contracts that were open as of December 31, 2024, a hypothetical 20% adverse movement in foreign currency exchange rates compared with the U.S. dollar relative to exchange rates as of December 31, 2024, would have resulted in a reduction in fair value of these contracts of approximately \$1.3 billion on this date and in the ensuing year, a reduction in income of \$700 million. The analysis does not consider the impact that hypothetical changes in foreign currency exchange rates would have on anticipated transactions that these foreign-currency-sensitive instruments were designed to offset.

As of December 31, 2025 and 2024, we had open, short-duration, foreign currency forward contracts that mature in one month or less, that had aggregate notional amounts of \$240 million and \$148 million, respectively, and that hedged fluctuations of certain assets and liabilities denominated in foreign currencies but were not designated as hedges for accounting purposes. These contracts had no material net unrealized gains or losses as of December 31, 2025 and 2024. With regard to these foreign currency forward contracts that were open as of December 31, 2025 and 2024, a hypothetical 5% adverse movement in foreign currency exchange rates compared with the U.S. dollar relative to exchange rates on these dates would not have a material effect on the fair values of these contracts or related income in the respective ensuing years. The analysis does not consider the impact that hypothetical changes in foreign currency exchange rates would have on assets and liabilities that these foreign-currency-sensitive instruments were designed to offset.

Market-price-sensitive financial instruments

As of December 31, 2025 and 2024, we were exposed to price risk on equity securities included in our portfolio of investments, which were acquired primarily for the promotion of business and strategic objectives. These investments include our investment in BeOne, as well as other publicly and privately held small-capitalization stocks and limited partnerships that invest in early-stage biotechnology companies. A 20% decrease in the aggregate value of our equity investment portfolio as of December 31, 2025 and 2024, would result in losses in fair value of approximately \$1.4 billion and \$950 million, respectively.

Counterparty credit risks

Our financial instruments, including derivatives, are subject to counterparty credit risk, which we consider as part of the overall fair value measurement. Our financial risk management policy limits derivative transactions by requiring that transactions be made only with institutions with minimum credit ratings of A- or equivalent by S&P, Moody's or Fitch; and it places exposure limits on the amount with any individual counterparty. In addition, we have an investment policy that limits

investments to certain types of debt and money market instruments issued by institutions with investment-grade credit ratings and places restriction on maturities and concentrations by asset class and issuer.

Item 8. FINANCIAL STATEMENTS AND SUPPLEMENTARY DATA

The information required by this item is incorporated herein by reference to the financial statements and schedule listed in Item 15(a)1 and (a)2 of Part IV and included in this Annual Report on Form 10-K.

Item 9. CHANGES IN AND DISAGREEMENTS WITH ACCOUNTANTS ON ACCOUNTING AND FINANCIAL DISCLOSURE

None.

Item 9A. CONTROLS AND PROCEDURES

We maintain “disclosure controls and procedures,” as defined in Rule 13a-15(e) under the Securities Exchange Act of 1934, that are designed to ensure that information required to be disclosed in Amgen’s Exchange Act reports is recorded, processed, summarized and reported within the time periods specified in the SEC’s rules and forms. These controls and procedures are also designed to ensure that such information is accumulated and communicated to Amgen’s management, including its Chief Executive Officer and Chief Financial Officer, as appropriate, to allow timely decisions regarding required disclosures. In designing and evaluating the disclosure controls and procedures, Amgen’s management recognized that any controls and procedures, no matter how well designed and operated, can provide only reasonable assurance of achieving the desired control objectives, and in reaching a reasonable level of assurance, Amgen’s management is required to apply its judgment in evaluating the cost-benefit relationship of possible controls and procedures. We have carried out an evaluation under the supervision and with the participation of our management, including Amgen’s Chief Executive Officer and Chief Financial Officer, of the effectiveness of the design and operation of Amgen’s disclosure controls and procedures. Based upon their evaluation and subject to the foregoing, our Chief Executive Officer and Chief Financial Officer concluded that our disclosure controls and procedures were effective as of December 31, 2025.

Management determined that as of December 31, 2025, there were no changes in our internal control over financial reporting that occurred during the fiscal quarter then ended that have materially affected or are reasonably likely to materially affect our internal control over financial reporting.

Management’s Report on Internal Control over Financial Reporting

Management of the Company is responsible for establishing and maintaining adequate internal control over financial reporting, as defined in Rule 13a-15(f) under the Securities Exchange Act of 1934. The Company’s internal control over financial reporting is designed to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with GAAP in the United States. However, all internal control systems, no matter how well designed, have inherent limitations. Therefore, even those systems determined to be effective can provide only reasonable assurance with respect to financial statement preparation and reporting.

Management assessed the effectiveness of the Company’s internal control over financial reporting as of December 31, 2025. In making this assessment, management used the criteria set forth by the COSO in Internal Control—Integrated Framework (2013 framework). Based on our assessment, management believes that the Company maintained effective internal control over financial reporting as of December 31, 2025, based on the COSO criteria.

The effectiveness of the Company’s internal control over financial reporting has been audited by Ernst & Young LLP, an independent registered public accounting firm, as stated in their attestation report appearing below, which expresses an unqualified opinion on the effectiveness of the Company’s internal control over financial reporting as of December 31, 2025.

Report of Independent Registered Public Accounting Firm

To the Stockholders and the Board of Directors of Amgen Inc.

Opinion on Internal Control Over Financial Reporting

We have audited Amgen Inc.'s internal control over financial reporting as of December 31, 2025, based on criteria established in Internal Control—Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission (2013 framework) (the COSO criteria). In our opinion, Amgen Inc. (the Company) maintained, in all material respects, effective internal control over financial reporting as of December 31, 2025, based on the COSO criteria.

We also have audited, in accordance with the standards of the Public Company Accounting Oversight Board (United States) (PCAOB), the consolidated balance sheets of the Company as of December 31, 2025 and 2024, the related consolidated statements of income, comprehensive income, stockholders' equity and cash flows for each of the three years in the period ended December 31, 2025, and the related notes and the financial statement schedule listed in the Index at Item 15(a)2 and our report dated February 13, 2026 expressed an unqualified opinion thereon.

Basis for Opinion

The Company's management is responsible for maintaining effective internal control over financial reporting and for its assessment of the effectiveness of internal control over financial reporting included in the accompanying Management's Report on Internal Control over Financial Reporting. Our responsibility is to express an opinion on the Company's internal control over financial reporting based on our audit. We are a public accounting firm registered with the PCAOB and are required to be independent with respect to the Company in accordance with the U.S. federal securities laws and the applicable rules and regulations of the Securities and Exchange Commission and the PCAOB.

We conducted our audit in accordance with the standards of the PCAOB. Those standards require that we plan and perform the audit to obtain reasonable assurance about whether effective internal control over financial reporting was maintained in all material respects.

Our audit included obtaining an understanding of internal control over financial reporting, assessing the risk that a material weakness exists, testing and evaluating the design and operating effectiveness of internal control based on the assessed risk, and performing such other procedures as we considered necessary in the circumstances. We believe that our audit provides a reasonable basis for our opinion.

Definition and Limitations of Internal Control Over Financial Reporting

A company's internal control over financial reporting is a process designed to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles. A company's internal control over financial reporting includes those policies and procedures that (1) pertain to the maintenance of records that, in reasonable detail, accurately and fairly reflect the transactions and dispositions of the assets of the company; (2) provide reasonable assurance that transactions are recorded as necessary to permit preparation of financial statements in accordance with generally accepted accounting principles, and that receipts and expenditures of the company are being made only in accordance with authorizations of management and directors of the company; and (3) provide reasonable assurance regarding prevention or timely detection of unauthorized acquisition, use, or disposition of the company's assets that could have a material effect on the financial statements.

Because of its inherent limitations, internal control over financial reporting may not prevent or detect misstatements. Also, projections of any evaluation of effectiveness to future periods are subject to the risk that controls may become inadequate because of changes in conditions, or that the degree of compliance with the policies or procedures may deteriorate.

/s/ Ernst & Young LLP

Los Angeles, California
February 13, 2026

Item 9B. OTHER INFORMATION

Rule 10b5-1 trading arrangements

During the three months ended December 31, 2025, none of our directors or officers (as defined in Rule 16a-1(f) of the Exchange Act) adopted or terminated any “Rule 10b5-1 trading arrangement” or “non-Rule 10b5-1 trading arrangement,” as each term is defined in Item 408 of Regulation S-K.

Item 9C. DISCLOSURE REGARDING FOREIGN JURISDICTIONS THAT PREVENT INSPECTIONS

Not applicable.

PART III

Item 10. DIRECTORS, EXECUTIVE OFFICERS AND CORPORATE GOVERNANCE

Information about our Directors is incorporated by reference from the section entitled ITEM 1—ELECTION OF DIRECTORS in our Proxy Statement for the 2026 Annual Meeting of Stockholders to be filed with the SEC within 120 days of December 31, 2025 (the Proxy Statement). Information about the procedures by which stockholders may recommend nominees for the Board of Directors is incorporated by reference from APPENDIX A—AMGEN INC. BOARD OF DIRECTORS GUIDELINES FOR DIRECTOR QUALIFICATIONS AND EVALUATIONS and OTHER MATTERS—Stockholder Proposals for the 2027 Annual Meeting in our Proxy Statement. Information about our Audit Committee, members of the committee and our Audit Committee financial experts is incorporated by reference from the section entitled CORPORATE GOVERNANCE—Audit Committee in our Proxy Statement. Information about our insider trading policies and procedures is incorporated by reference from the section entitled COMPENSATION DISCUSSION AND ANALYSIS—COMPENSATION POLICIES AND PRACTICES—Insider Trading Policies and Procedures in our Proxy Statement. A copy of our Insider Trading Policy and our securities transactions pre-clearance practices and procedures are incorporated by reference to this Annual Report on Form 10-K as Exhibits 19.1 and 19.2, respectively. Information about our executive officers is contained in the discussion entitled Part I, Item 1. Business—Information about our Executive Officers.

Code of Ethics

We maintain a Code of Ethics for the Chief Executive Officer and Senior Financial Officers applicable to our principal executive officer, principal financial officer, principal accounting officer or controller and other persons performing similar functions. To view this code of ethics free of charge, please visit our website at www.amgen.com. (The website address is not intended to function as a hyperlink, and the information contained in our website is not intended to be a part of this filing.) We intend to satisfy the disclosure requirements under Item 5.05 of Form 8-K regarding an amendment to or a waiver from a provision of this code of ethics, if any, by posting such information on our website as set forth above.

Item 11. EXECUTIVE COMPENSATION

Information about director and executive compensation is incorporated by reference from the sections entitled COMPENSATION DISCUSSION AND ANALYSIS, EXECUTIVE COMPENSATION TABLES, DIRECTOR COMPENSATION and OTHER COMPENSATION MATTERS in our Proxy Statement. Information about compensation committee matters is incorporated by reference from the sections entitled CORPORATE GOVERNANCE—Compensation and Management Development Committee and COMPENSATION COMMITTEE REPORT in our Proxy Statement.

Item 12. SECURITY OWNERSHIP OF CERTAIN BENEFICIAL OWNERS AND MANAGEMENT AND RELATED STOCKHOLDER MATTERS

Securities Authorized for Issuance Under Existing Equity Compensation Plans

The following table sets forth certain information as of December 31, 2025, concerning the shares of our common stock that may be issued under any form of award granted under our equity compensation plans in effect as of December 31, 2025, including upon the exercise of options, upon the vesting of awards of RSUs or when performance units are earned and related dividend equivalents have been granted.

Plan category	(a) Number of securities to be issued upon exercise of outstanding options and rights	(b) Weighted-average exercise price of outstanding options and rights	(c) Number of securities remaining available for future issuance under equity compensation plans (excluding securities reflected in column (a))
Equity compensation plans approved by Amgen security holders:			
Amended and Restated 2009 Equity Incentive Plan ⁽¹⁾	11,098,368	\$ 234.45	35,004,385
Amended and Restated 1991 Equity Incentive Plan ⁽²⁾	100		
Amended and Restated Employee Stock Purchase Plan			3,813,679
Total approved plans	11,098,468	\$ 234.45	38,818,064
Equity compensation plan not approved by Amgen security holders:			
Amgen Profit Sharing Plan for Employees in Ireland ⁽³⁾			150,740
Horizon stock plans ⁽⁴⁾	132,521		
Total unapproved plans	132,521	—	150,740
Total all plans	11,230,989	\$ 234.45	38,968,804

⁽¹⁾ The Amended 2009 Plan employs a fungible share-counting formula for determining the number of shares available for issuance under the plan. In accordance with this formula, each option or stock appreciation right counts as one share, while each RSU, performance unit or dividend equivalent counts as 1.9 shares. The number under column (a) represents the actual number of shares issuable under our outstanding awards without giving effect to the fungible share-counting formula. The number under column (c) represents the number of shares available for issuance under this plan based on each such available share counting as one share. Commencing with the grants made in April 2012, RSUs and performance units accrue dividend equivalents that are payable in shares only to the extent and when the underlying RSUs vest or underlying performance units have been earned and the related shares are issued to the grantee. The performance units granted under this plan are earned based on the accomplishment of specified performance goals at the end of their respective three-year performance periods; the number of performance units granted represent target performance, and the maximum number of units that could be earned based on our performance is 200% of the performance units granted in 2023, 2024 and 2025.

As of December 31, 2025, the number of outstanding awards under column (a) includes (i) 6,064,625 shares issuable upon the exercise of outstanding options with a weighted-average exercise price of \$234.45; (ii) 3,797,669 shares issuable upon the vesting of outstanding RSUs, including 221,796 related dividend equivalents; and (iii) 1,236,074 shares subject to outstanding 2023, 2024 and 2025 performance units, including 74,616 related dividend equivalents. The weighted-average exercise price shown in column (b) is for the outstanding options only. The number of available shares under column (c) represents the number of shares that remain available for future issuance under this plan as of December 31, 2025, employing the fungible share formula and presumes the issuance of target shares under the performance units granted in 2023, 2024 and 2025 and related dividend equivalents. The numbers under columns (a) and (c) do not give effect to the additional shares that could be issuable in the event that above target performance on the performance goals under these outstanding performance units is achieved. Maximum performance under these goals could result in 200% of target shares being awarded for performance units granted in 2023, 2024 and 2025.

⁽²⁾ This plan has terminated as to future grants. The number under column (a) with respect to this plan includes 100 shares issuable upon the settlement of deferred RSUs.

⁽³⁾ The Profit Sharing Plan was approved by the Board of Directors on July 28, 2011. The Profit Sharing Plan permits eligible employees of the Company's subsidiaries located in Ireland who participate in the Profit Sharing Plan to apply a

portion of their qualifying bonus and salary to the purchase of the Company's common stock on the open market at the market price by a third-party trustee as described in the Profit Sharing Plan.

- ⁽⁴⁾ The Horizon Therapeutics Public Limited Company Amended and Restated 2014 Equity Incentive Plan, the Horizon Therapeutics Public Limited Company Amended and Restated 2020 Equity Incentive Plan and 2020 Restricted Stock Unit Award Sub-Plan and the Horizon Therapeutics Public Limited Company Amended and Restated 2018 Equity Incentive Plan and 2018 Restricted Stock Unit Award Sub-Plan (collectively, the "Horizon stock plans") were acquired on October 6, 2023, pursuant to our acquisition of Horizon. In connection with the closing of the Horizon acquisition and pursuant to its terms, outstanding RSUs issued under the Horizon stock plans were converted into Amgen RSUs, and these plans terminated as to future grants on October 6, 2023.

Security Ownership of Directors and Executive Officers and Certain Beneficial Owners

Information about security ownership of certain beneficial owners and management is incorporated by reference from the sections entitled SECURITY OWNERSHIP OF DIRECTORS AND EXECUTIVE OFFICERS and SECURITY OWNERSHIP OF CERTAIN BENEFICIAL OWNERS in our Proxy Statement.

Item 13. CERTAIN RELATIONSHIPS AND RELATED TRANSACTIONS AND DIRECTOR INDEPENDENCE

Information about certain relationships and related transactions and director independence is incorporated by reference from the sections entitled CERTAIN RELATIONSHIPS AND RELATED TRANSACTIONS and CORPORATE GOVERNANCE—Director Independence in our Proxy Statement.

Item 14. PRINCIPAL ACCOUNTANT FEES AND SERVICES

Information about the fees for professional services rendered by our independent registered public accountants is incorporated by reference from the section entitled AUDIT MATTERS—Independent Registered Public Accountants in our Proxy Statement.

PART IV

Item 15. EXHIBITS AND FINANCIAL STATEMENT SCHEDULES

(a)1. Index to Financial Statements

The following Consolidated Financial Statements are included herein:

	Page number
Report of Independent Registered Public Accounting Firm (PCAOB ID: 42)	F-1
Consolidated Statements of Income for each of the three years in the period ended December 31, 2025	F-4
Consolidated Statements of Comprehensive Income for each of the three years in the period ended December 31, 2025	F-5
Consolidated Balance Sheets as of December 31, 2025 and 2024	F-6
Consolidated Statements of Stockholders' Equity for each of the three years in the period ended December 31, 2025	F-7
Consolidated Statements of Cash Flows for each of the three years in the period ended December 31, 2025	F-8
Notes to Consolidated Financial Statements	F-9

(a)2. Index to Financial Statement Schedules

The following Schedule is filed as part of this Annual Report on Form 10-K:

	Page number
Schedule II. Valuation and Qualifying Accounts	F-61

All other schedules are omitted because they are not applicable, not required or because the required information is included in the consolidated financial statements or notes thereto.

(a)3. Exhibits

Exhibit No.	Description
3.1	Restated Certificate of Incorporation of Amgen Inc. (As Restated March 6, 2013.) (Filed as an exhibit to Form 10-Q for the quarter ended March 31, 2013 on May 3, 2013 and incorporated herein by reference.)
3.2	Amended and Restated Bylaws of Amgen Inc. (As Amended and Restated February 15, 2016.) (Filed as an exhibit to Form 8-K on February 17, 2016 and incorporated herein by reference.)
4.1	Form of stock certificate for the common stock, par value \$.0001 of the Company. (Filed as an exhibit to Form 10-Q for the quarter ended March 31, 1997 on May 14, 1997 and incorporated herein by reference.)
4.2	Form of Indenture, dated January 1, 1992. (Filed as an exhibit to Form S-3 Registration Statement filed on December 19, 1991 and incorporated herein by reference.)
4.3	Agreement of Resignation, Appointment and Acceptance dated February 15, 2008. (Filed as an exhibit to Form 10-K for the year ended December 31, 2007 on February 28, 2008 and incorporated herein by reference.)
4.4	First Supplemental Indenture, dated February 26, 1997. (Filed as an exhibit to Form 8-K on March 14, 1997 and incorporated herein by reference.)
4.5	8-1/8% Debentures due April 1, 2097. (Filed as an exhibit to Form 8-K on April 8, 1997 and incorporated herein by reference.)
4.6	Officer's Certificate of Amgen Inc., dated April 8, 1997, establishing a series of securities entitled "8 1/8% Debentures due April 1, 2097." (Filed as an exhibit to Form 8-K on April 8, 1997 and incorporated herein by reference.)

Exhibit No.	Description
4.7	<u>Indenture, dated August 4, 2003</u> , (Filed as an exhibit to Form S-3 Registration Statement on August 4, 2003 and incorporated herein by reference.)
4.8	<u>Corporate Commercial Paper - Master Note between and among Amgen Inc., as Issuer, Cede & Co., as Nominee of The Depository Trust Company, and Citibank, N.A., as Paying Agent</u> , (Filed as an exhibit to Form 10-Q for the quarter ended March 31, 1998 on May 13, 1998 and incorporated herein by reference.)
4.9	<u>Officers' Certificate of Amgen Inc., dated May 30, 2007, including form of the Company's 6.375% Senior Notes due 2037</u> , (Filed as an exhibit to Form 8-K on May 30, 2007 and incorporated herein by reference.)
4.10	<u>Officers' Certificate of Amgen Inc., dated May 23, 2008, including form of the Company's 6.90% Senior Notes due 2038</u> , (Filed as exhibit to Form 8-K on May 23, 2008 and incorporated herein by reference.)
4.11	<u>Officers' Certificate of Amgen Inc., dated January 16, 2009, including form of the Company's 6.40% Senior Notes due 2039</u> , (Filed as exhibit to Form 8-K on January 16, 2009 and incorporated herein by reference.)
4.12	<u>Officers' Certificate of Amgen Inc., dated March 12, 2010, including form of the Company's 5.75% Senior Notes due 2040</u> , (Filed as exhibit to Form 8-K on March 12, 2010 and incorporated herein by reference.)
4.13	<u>Officers' Certificate of Amgen Inc., dated September 16, 2010, including form of the Company's 4.95% Senior Notes due 2041</u> , (Filed as an exhibit to Form 8-K on September 17, 2010 and incorporated herein by reference.)
4.14	<u>Officers' Certificate of Amgen Inc., dated June 30, 2011, including form of the Company's 5.65% Senior Notes due 2042</u> , (Filed as an exhibit to Form 8-K on June 30, 2011 and incorporated herein by reference.)
4.15	<u>Officers' Certificate of Amgen Inc., dated November 10, 2011, including form of the Company's 5.15% Senior Notes due 2041</u> , (Filed as an exhibit to Form 8-K on November 10, 2011 and incorporated herein by reference.)
4.16	<u>Officers' Certificate of Amgen Inc., dated December 5, 2011, including form of the Company's 5.50% Senior Notes due 2026</u> , (Filed as an exhibit to Form 8-K on December 5, 2011 and incorporated herein by reference.)
4.17	<u>Officers' Certificate of Amgen Inc., dated May 15, 2012, including form of the Company's 5.375% Senior Notes due 2043</u> , (Filed as an exhibit to Form 8-K on May 15, 2012 and incorporated herein by reference.)
4.18	<u>Officers' Certificate of Amgen Inc., dated September 13, 2012, including form of the Company's 4.000% Senior Notes due 2029</u> , (Filed as an exhibit to Form 8-K on September 13, 2012 and incorporated herein by reference.)
4.19	<u>Indenture, dated May 22, 2014, between Amgen Inc. and The Bank of New York Mellon Trust Company, N.A., as Trustee</u> , (Filed as an exhibit to Form 8-K on May 22, 2014 and incorporated herein by reference.)
4.20	<u>Officer's Certificate of Amgen Inc., dated May 1, 2015, including forms of the Company's 3.125% Senior Notes due 2025 and 4.400% Senior Notes due 2045</u> , (Filed as an exhibit on Form 8-K on May 1, 2015 and incorporated herein by reference.)
4.21	<u>Officer's Certificate of Amgen Inc., dated as of February 25, 2016, including form of the Company's 2.000% Senior Notes due 2026</u> , (Filed as an exhibit on Form 8-K on February 26, 2016 and incorporated herein by reference.)
4.22	<u>Officer's Certificate of Amgen Inc., dated as of June 14, 2016, including forms of the Company's 4.563% Senior Notes due 2048 and 4.663% Senior Notes due 2051</u> , (Filed as an exhibit to Form 8-K on June 14, 2016 and incorporated herein by reference.)
4.23	<u>Officer's Certificate of Amgen Inc., dated as of August 19, 2016, including forms of the Company's 2.600% Senior Notes due 2026</u> , (Filed as an exhibit to Form 8-K on August 19, 2016 and incorporated herein by reference.)
4.24	<u>Officer's Certificate of Amgen Inc., dated as of November 2, 2017, including in the form of the Company's 3.200% Senior Notes due 2027</u> , (Filed as an exhibit to Form 8-K on November 2, 2017 and incorporated herein by reference.)
4.25	<u>Officer's Certificate of Amgen Inc., dated as of February 21, 2020, including forms of the Company's 1.900% Senior Notes due 2025, 2.200% Senior Notes due 2027, 2.450% Senior Notes due 2030, 3.150% Senior Notes due 2040 and 3.375% Senior Notes due 2050</u> , (Filed as an exhibit to Form 8-K on February 21, 2020 and incorporated herein by reference.)

Exhibit No.	Description
4.26	<u>Officer's Certificate of Amgen Inc., dated as of May 6, 2020, including form of the Company's 2.300% Senior Notes due 2031.</u> (Filed as an exhibit to Form 8-K on May 6, 2020 and incorporated herein by reference.)
4.27	<u>Officer's Certificate of Amgen Inc., dated as of August 17, 2020, including forms of the Company's 2.770% Senior Notes due 2053.</u> (Filed as an exhibit to Form 8-K on August 18, 2020 and incorporated herein by reference.)
4.28	<u>Officer's Certificate of Amgen Inc., dated as of August 9, 2021, including forms of the Company's 1.650% Senior Notes due 2028, 2.000% Senior Notes due 2032, 2.800% Senior Notes due 2041 and 3.000% Senior Notes due 2052.</u> (Filed as an exhibit to Form 8-K on August 9, 2021 and incorporated herein by reference.)
4.29	<u>Officer's Certificate of Amgen Inc., dated as of February 22, 2022, including forms of the Company's 3.000% Senior Notes due 2029, 3.350% Senior Notes due 2032, 4.200% Senior Notes due 2052 and 4.400% Senior Notes due 2062.</u> (Filed as an exhibit to Form 8-K on February 22, 2022 and incorporated herein by reference.)
4.30	<u>Officer's Certificate of Amgen Inc., dated as of August 18, 2022, including forms of the Company's 4.050% Senior Notes due 2029, 4.200% Senior Notes due 2033 and 4.875% Senior Notes due 2053.</u> (Filed as an exhibit to Form 8-K on August 18, 2022 and incorporated herein by reference.)
4.31	<u>Officer's Certificate of the Company, dated as of March 2, 2023, including forms of the Company's 5.250% Senior Notes due 2025, 5.507% Senior Notes due 2026, 5.150% Senior Notes due 2028, 5.250% Senior Notes due 2030, 5.250% Senior Notes due 2033, 5.600% Senior Notes due 2043, 5.650% Senior Notes due 2053 and 5.750% Senior Notes due 2063.</u> (Filed as an exhibit to Form 8-K on March 2, 2023 and incorporated herein by reference.)
4.32*	<u>Description of Amgen Inc.'s Securities Registered Pursuant to Section 12 of the Securities Exchange Act of 1934.</u>
10.1+	<u>Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan.</u> (Filed as Appendix C to the Definitive Proxy Statement on Schedule 14A on April 17, 2024 and incorporated herein by reference.)
10.2*+	<u>Form of Grant of Stock Option Agreement for the Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan.</u> (As Amended and Restated on December 8, 2025.)
10.3*+	<u>Form of Restricted Stock Unit Agreement for the Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan.</u> (As Amended and Restated on December 8, 2025.)
10.4+	<u>Amgen Inc. 2009 Performance Award Program.</u> (As Amended and Restated on May 31, 2024.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2024 on August 7, 2024 and incorporated herein by reference.)
10.5*+	<u>Form of Performance Unit Agreement for the Amgen Inc. 2009 Performance Award Program.</u> (As Amended and Restated on December 8, 2025.)
10.6+	<u>Amgen Inc. 2009 Director Equity Incentive Program.</u> (As Amended and Restated on May 31, 2024.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2024 on August 7, 2024 and incorporated herein by reference.)
10.7+	<u>Form of Restricted Stock Unit Agreement for the Amgen Inc. 2009 Director Equity Incentive Program.</u> (As Amended and Restated on May 31, 2024.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2024 on August 7, 2024 and incorporated herein by reference.)
10.8+	<u>Form of Cash-Settled Restricted Stock Unit Agreement for the Amgen Inc. 2009 Director Equity Incentive Program.</u> (As Amended and Restated on May 31, 2024.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2024 on August 7, 2024 and incorporated herein by reference.)
10.9+	<u>Amgen Inc. Supplemental Retirement Plan. (As Amended and Restated effective October 16, 2013.)</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2013 on February 24, 2014 and incorporated herein by reference.)
10.9.1+	<u>First Amendment to the Amgen Inc. Supplemental Retirement Plan, effective October 14, 2016.</u> (Filed as an exhibit to Form 10-Q for the quarter ended September 30, 2016 on October 28, 2016 and incorporated herein by reference.)

Exhibit No.	Description
10.9.2+	<u>Second Amendment to the Amgen Inc. Supplemental Retirement Plan, effective October 23, 2019</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2019 on February 12, 2020 and incorporated herein by reference.)
10.9.3+	<u>Third Amendment to the Amgen Inc. Supplemental Retirement Plan, effective October 20, 2021</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2021 on February 16, 2022 and incorporated herein by reference.)
10.9.4+	<u>Fourth Amendment to the Amgen Inc. Supplemental Retirement Plan, effective October 20, 2022</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2022 on February 9, 2023 and incorporated herein by reference.)
10.9.5+	<u>Fifth Amendment to the Amgen Inc. Supplemental Retirement Plan, effective January 1, 2024</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2023 on February 14, 2024 and incorporated herein by reference.)
10.10+	<u>Amended and Restated Amgen Change of Control Severance Plan. (As Amended and Restated effective December 9, 2010 and subsequently amended effective March 2, 2011.)</u> (Filed as an exhibit to Form 10-Q for the quarter ended March 31, 2011 on May 10, 2011 and incorporated herein by reference.)
10.11+	<u>Amgen Inc. Executive Incentive Plan</u> (As Amended and Restated effective January 1, 2022.) (Filed as an exhibit to Form 10-Q for the quarter ended March 31, 2022 on April 28, 2022 and incorporated herein by reference.)
10.12+	<u>Amgen Nonqualified Deferred Compensation Plan. (As Amended and Restated effective October 16, 2013.)</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2013 on February 24, 2014 and incorporated herein by reference.)
10.12.1+	<u>First Amendment to the Amgen Nonqualified Deferred Compensation Plan, effective October 14, 2016</u> (Filed as an exhibit to Form 10-Q for the quarter ended September 30, 2016 on October 28, 2016 and incorporated herein by reference.)
10.12.2+	<u>Second Amendment to the Amgen Nonqualified Deferred Compensation Plan, effective January 1, 2020</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2019 on February 12, 2020 and incorporated herein by reference.)
10.12.3+	<u>Third Amendment to the Amgen Nonqualified Deferred Compensation Plan, effective January 1, 2022</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2021 on February 16, 2022 and incorporated herein by reference.)
10.12.4+	<u>Fourth Amendment to the Amgen Nonqualified Deferred Compensation Plan, effective January 1, 2024</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2023 on February 14, 2024 and incorporated herein by reference.)
10.13+	<u>Aircraft Time Sharing Agreement, dated December 3, 2021, by and between Amgen Inc. and Robert A. Bradway</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2021 on February 16, 2022 and incorporated herein by reference.)
10.14+	<u>Agreement between Amgen Inc. and James Bradner, dated December 13, 2023</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2023 on February 14, 2024 and incorporated herein by reference.)
10.15	<u>Term Loan Credit Agreement, dated as of December 22, 2022, by and among Amgen Inc., Citibank, N.A., as administrative agent, Bank of America, N.A., as syndication agent, Citibank, N.A., Bank of America, N.A., Goldman Sachs Bank USA and Mizuho Bank, Ltd., as lead arrangers and book runners, Goldman Sachs Bank USA and Mizuho Bank, Ltd. as documentation agents, and the other banks party thereto</u> (Filed as an exhibit to Form 8-K on December 22, 2022 and incorporated herein by reference.)
10.16	<u>Third Amended and Restated Credit Agreement, dated as of March 9, 2023, among Amgen Inc., the Banks therein named, Citibank, N.A., as Administrative Agent, and JPMorgan Chase Bank, N.A., as Syndication Agent</u> (Filed as an exhibit to Form 8-K on March 9, 2023 and incorporated herein by reference.)

Exhibit No.	Description
10.17	<u>Collaboration and License Agreement between Amgen Inc. and Celltech R&D Limited dated May 10, 2002</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) <u>and Amendment No. 1, effective June 9, 2003, to Collaboration and License Agreement between Amgen Inc. and Celltech R&D Limited</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-K for the year ended December 31, 2023 on February 14, 2024 and incorporated herein by reference.)
10.17.1	<u>Amendment No. 2 to Collaboration and License Agreement, effective November 14, 2016, between Amgen Inc. and Celltech R&D Limited</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-K for the year ended December 31, 2023 on February 14, 2024 and incorporated herein by reference.)
10.18	<u>Letter Agreement, dated June 25, 2019, by and between Amgen Inc. and UCB Celltech (portions of the exhibit have been omitted because they are both (i) not material and (ii) would be competitively harmful if publicly disclosed)</u> (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2019 on July 31, 2019 and incorporated herein by reference.)
10.19	<u>Collaboration Agreement, dated October 31, 2019, by and between Amgen Inc. and BeiGene Switzerland GmbH, a wholly-owned subsidiary of BeiGene, Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) would be competitively harmful if publicly disclosed.) (Filed as an exhibit to Form 10-K for the year ended December 31, 2019 on February 12, 2020 and incorporated herein by reference.)
10.19.1	<u>First Amendment to Collaboration Agreement, dated April 20, 2022, by and between Amgen Inc. and BeiGene Switzerland GmbH, and BeiGene, Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2022 on August 5, 2022 and incorporated herein by reference.)
10.19.2	<u>Second Amendment to Collaboration Agreement, entered into as of February 26, 2023, by and between Amgen Inc. and BeiGene Switzerland GmbH, and BeiGene, Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended March 31, 2023 on April 28, 2023 and incorporated herein by reference.)
10.19.3	<u>Letter Agreement, dated May 9, 2025, by and between Amgen Inc. and BeiGene Switzerland GmbH, a wholly owned subsidiary of BeiGene, Ltd.</u> (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2025 on August 5, 2025 and incorporated herein by reference.)
10.19.4	<u>Letter Agreement, dated August 11, 2025, by and between Amgen Inc. and BeOne Medicines I GmbH and BeOne Medicines Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended September 30, 2025 on November 5, 2025 and incorporated herein by reference.)
10.19.5*	<u>Letter Agreement, dated October 1, 2025, by and between Amgen Inc. and BeOne Medicines I GmbH and BeOne Medicines Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.)
10.19.6*	<u>Third Amendment to Collaboration Agreement, dated October 31, 2025, by and among Amgen Inc., BeOne Medicines I GmbH, and BeOne Medicines Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.)
10.19.7*	<u>Fourth Amendment to Collaboration Agreement, dated November 11, 2025, by and among Amgen Inc., BeOne Medicines I GmbH, and BeOne Medicines Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.)
10.20	<u>Guarantee, dated as of October 31, 2019, made by and among BeiGene, Ltd. and Amgen Inc.</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2019 on February 12, 2020 and incorporated herein by reference.)
10.21	<u>Share Purchase Agreement, dated October 31, 2019, by and between Amgen Inc. and BeiGene, Ltd.</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) would be competitively harmful if publicly disclosed.) (Filed as an exhibit to Schedule 13D on January 8, 2020 and incorporated herein by reference.)

Exhibit No.	Description
10.21.1	<u>Amendment No. 1 to Share Purchase Agreement, dated December 6, 2019, by and among BeiGene, Ltd. and Amgen Inc.</u> (Filed as an exhibit to Schedule 13D on January 8, 2020 and incorporated herein by reference.)
10.21.2	<u>Restated Amendment No. 2 to Share Purchase Agreement, dated September 24, 2020, by and among BeiGene, Ltd. and Amgen Inc.</u> (Filed as an exhibit to Form 10-Q for the quarter ended September 30, 2020 on October 29, 2020 and incorporated herein by reference.)
10.21.3	<u>Amendment No. 3 to Share Purchase Agreement, dated January 30, 2023, by and among BeiGene, Ltd. and Amgen Inc.</u> (Filed as an exhibit to Form 8-K on January 31, 2023 and incorporated herein by reference.)
10.22	<u>Collaboration Agreement dated March 30, 2012 by and between Amgen Inc. and AstraZeneca Collaboration Ventures, LLC, a wholly owned subsidiary of AstraZeneca Pharmaceuticals LP</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2022 on August 5, 2022 and incorporated herein by reference.)
10.22.1	<u>Amendment No. 1 to the Collaboration Agreement, dated October 1, 2014, by and among Amgen Inc., AstraZeneca Collaboration Ventures, LLC and AstraZeneca Pharmaceuticals LP</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2022 on August 5, 2022 and incorporated herein by reference.)
10.22.2	<u>Amendment Nos. 2 through 6 to the March 30, 2012 Collaboration Agreement between Amgen Inc. and AstraZeneca Collaboration Ventures, LLC, dated May 2 and 27 and October 2, 2016, January 31, 2018, and May 15, 2020, respectively</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) would be competitively harmful if publicly disclosed.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2020 on July 29, 2020 and incorporated herein by reference.)
10.22.3	<u>Amendment No. 7 to the Collaboration Agreement, dated December 17, 2020, by and between Amgen Inc. and AstraZeneca Collaboration Ventures, LLC</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) would be competitively harmful if publicly disclosed.) (Filed as an exhibit to Form 10-K for the year ended December 31, 2020 on February 9, 2021 and incorporated herein by reference.)
10.22.4	<u>Amendment No. 8 to the Collaboration Agreement, dated November 19, 2021, by and between Amgen Inc. and AstraZeneca Collaboration Ventures, LLC</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-K for the year ended December 31, 2021 on February 16, 2022 and incorporated herein by reference.)
10.22.5	<u>Letter Agreement Regarding the Collaboration Agreement, dated as of December 1, 2023, by and between Amgen Inc. and AstraZeneca Collaboration Ventures, LLC</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-K for the year ended December 31, 2023 on February 14, 2024 and incorporated herein by reference.)
10.22.6	<u>Amendment No. 9 to the Collaboration Agreement, dated May 20, 2025, by and between Amgen Inc. and AstraZeneca Collaboration Ventures, LLC</u> (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2025 on August 5, 2025 and incorporated herein by reference.)
10.23	<u>License and Collaboration Agreement, dated June 1, 2021, by and between Amgen Inc. and Kyowa Kirin Co., Ltd.</u> ² (portions of the exhibit have been omitted because they are both (i) not material and (ii) is the type of information that the Company treats as private or confidential.) (Filed as an exhibit to Form 10-Q for the quarter ended June 30, 2021 on August 4, 2021 and incorporated herein by reference.)
19.1	<u>Amgen Inc. Insider Trading Policy</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2024 on February 14, 2025 and incorporated herein by reference.)
19.2	<u>Amgen Inc. Securities Transactions Blackout and Pre-Clearance Practices and Procedures</u> (Filed as an exhibit to Form 10-K for the year ended December 31, 2024 on February 14, 2025 and incorporated herein by reference.)
21*	<u>Subsidiaries of the Company</u>

Exhibit No.	Description
23	Consent of the Independent Registered Public Accounting Firm. The consent is set forth on page 96 of this Annual Report on Form 10-K.
24	Power of Attorney. The Power of Attorney is set forth on page 97 of this Annual Report on Form 10-K.
31*	Rule 13a-14(a) Certifications.
32**	Section 1350 Certifications.
97	Policy Relating to Recovery of Erroneously Awarded Compensation. (Filed as an exhibit to Form 10-K for the year ended December 31, 2024 on February 14, 2025 and incorporated herein by reference.)
101.INS	Inline XBRL Instance Document - The instance document does not appear in the interactive data file because its XBRL tags are embedded within the Inline XBRL document.
101.SCH*	Inline XBRL Taxonomy Extension Schema Document.
101.CAL*	Inline XBRL Taxonomy Extension Calculation Linkbase Document.
101.DEF*	Inline XBRL Taxonomy Extension Definition Linkbase Document.
101.LAB*	Inline XBRL Taxonomy Extension Label Linkbase Document.
101.PRE*	Inline XBRL Taxonomy Extension Presentation Linkbase Document.
104	Cover Page Interactive Data File (formatted as Inline XBRL and contained in Exhibit 101).

* = filed herewith

** = furnished herewith and not “filed” for purposes of Section 18 of the Securities Exchange Act of 1934, as amended

+ = management contract or compensatory plan or arrangement

¹ In May 2025, BeiGene, Ltd. changed its name to BeOne Medicines Ltd., and BeiGene Switzerland GmbH changed its name to BeOne Medicines I GmbH.

² See Significant Developments for additional information regarding the termination of our collaboration agreement with Kyowa Kirin.

Item 16. FORM 10-K SUMMARY

Not applicable.

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this Annual Report to be signed on its behalf by the undersigned, thereunto duly authorized.

AMGEN INC.
(Registrant)

Date: February 13, 2026

By:

/s/ PETER H. GRIFFITH

Peter H. Griffith
Executive Vice President and Chief Financial Officer
(Principal Financial Officer)

CONSENT OF INDEPENDENT REGISTERED PUBLIC ACCOUNTING FIRM

We consent to the incorporation by reference in the following Registration Statements:

- Registration Statement (Form S-3 No. 333-269670) of Amgen Inc.,
- Registration Statements (Form S-8 Nos. 333-159377 and 333-280155) pertaining to the Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan,
- Registration Statement (Form S-8 No. 33-39183) pertaining to the Amgen Inc. Amended and Restated Employee Stock Purchase Plan,
- Registration Statements (Form S-8 Nos. 33-39104, 333-144581 and 333-216719) pertaining to the Amgen Retirement and Savings Plan,
- Registration Statements (Form S-8 Nos. 33-47605, 333-144580 and 333-216715) pertaining to The Retirement and Savings Plan for Amgen Manufacturing Limited LLC (formerly known as The Retirement and Savings Plan for Amgen Manufacturing, Limited),
- Registration Statements (Form S-8 Nos. 333-81284, 333-177868, 333-216723 and 333-260723) pertaining to the Amgen Nonqualified Deferred Compensation Plan,
- Registration Statements (Form S-8 Nos. 333-176240 and 333-260724) pertaining to the Amgen Profit Sharing Plan for Employees in Ireland, and
- Registration Statement (Form S-8 No. 333-274900) pertaining to the Horizon Therapeutics Public Limited Company Amended and Restated 2014 Equity Incentive Plan, Horizon Therapeutics Public Limited Company Amended and Restated 2018 Equity Incentive Plan and 2018 Restricted Stock Unit Award Sub-Plan, and Horizon Therapeutics Public Limited Company Amended and Restated 2020 Equity Incentive Plan and 2020 Restricted Stock Unit Award Sub-Plan;

of our reports dated February 13, 2026, with respect to the consolidated financial statements of Amgen Inc. and the effectiveness of internal control over financial reporting of Amgen Inc. included in this Annual Report (Form 10-K) of Amgen Inc. for the year ended December 31, 2025.

/s/ Ernst & Young LLP

Los Angeles, California
February 13, 2026

POWER OF ATTORNEY

KNOW ALL MEN AND WOMEN BY THESE PRESENTS, that each person whose signature appears below constitutes and appoints Robert A. Bradway, Peter H. Griffith and Jonathan P. Graham, or any of them, his or her attorney-in-fact, each with the power of substitution and re-substitution, for him or her in any and all capacities, to sign any amendments to this Report, and to file the same, with exhibits thereto and other documents in connection therewith, with the Securities and Exchange Commission, hereby ratifying and confirming all that each of said attorneys-in-fact, or his or her substitute or substitutes, may do or cause to be done by virtue hereof.

Pursuant to the requirements of the Securities Exchange Act of 1934, this report has been signed below by the following persons on behalf of the registrant and in the capacities and on the dates indicated:

Signature	Title	Date
/S/ ROBERT A. BRADWAY Robert A. Bradway	Chairman of the Board, Chief Executive Officer and President, and Director (Principal Executive Officer)	2/13/2026
/S/ PETER H. GRIFFITH Peter H. Griffith	Executive Vice President and Chief Financial Officer (Principal Financial Officer)	2/13/2026
/S/ MATTHEW C. BUSCH Matthew C. Busch	Vice President, Finance and Chief Accounting Officer (Principal Accounting Officer)	2/13/2026
/S/ WANDA M. AUSTIN Wanda M. Austin	Director	2/13/2026
/S/ MICHAEL V. DRAKE Michael V. Drake	Director	2/13/2026
/S/ BRIAN J. DRUKER Brian J. Druker	Director	2/13/2026
/S/ ROBERT A. ECKERT Robert A. Eckert	Director	2/13/2026
/S/ GREG C. GARLAND Greg C. Garland	Director	2/13/2026
/S/ CHARLES M. HOLLEY, JR. Charles M. Holley, Jr.	Director	2/13/2026
/S/ S. OMAR ISHRAK S. Omar Ishrak	Director	2/13/2026
/S/ TYLER JACKS Tyler Jacks	Director	2/13/2026
/S/ MARY E. KLOTMAN Mary E. Klotman	Director	2/13/2026
/S/ ELLEN J. KULLMAN Ellen J. Kullman	Director	2/13/2026
/S/ AMY E. MILES Amy E. Miles	Director	2/13/2026

Report of Independent Registered Public Accounting Firm

To the Stockholders and the Board of Directors of Amgen Inc.

Opinion on the Financial Statements

We have audited the accompanying consolidated balance sheets of Amgen Inc. (the Company) as of December 31, 2025 and 2024, the related consolidated statements of income, comprehensive income, stockholders' equity and cash flows for each of the three years in the period ended December 31, 2025, and the related notes and financial statement schedule listed in the Index at Item 15(a)2 (collectively referred to as the "consolidated financial statements"). In our opinion, the consolidated financial statements present fairly, in all material respects, the financial position of the Company at December 31, 2025 and 2024, and the results of its operations and its cash flows for each of the three years in the period ended December 31, 2025, in conformity with U.S. generally accepted accounting principles.

We also have audited, in accordance with the standards of the Public Company Accounting Oversight Board (United States) (PCAOB), the Company's internal control over financial reporting as of December 31, 2025, based on criteria established in Internal Control—Integrated Framework issued by the Committee of Sponsoring Organizations of the Treadway Commission (2013 framework) and our report dated February 13, 2026 expressed an unqualified opinion thereon.

Basis for Opinion

These financial statements are the responsibility of the Company's management. Our responsibility is to express an opinion on the Company's financial statements based on our audits. We are a public accounting firm registered with the PCAOB and are required to be independent with respect to the Company in accordance with the U.S. federal securities laws and the applicable rules and regulations of the Securities and Exchange Commission and the PCAOB.

We conducted our audits in accordance with the standards of the PCAOB. Those standards require that we plan and perform the audit to obtain reasonable assurance about whether the financial statements are free of material misstatement, whether due to error or fraud. Our audits included performing procedures to assess the risks of material misstatement of the financial statements, whether due to error or fraud, and performing procedures that respond to those risks. Such procedures included examining, on a test basis, evidence regarding the amounts and disclosures in the financial statements. Our audits also included evaluating the accounting principles used and significant estimates made by management, as well as evaluating the overall presentation of the financial statements. We believe that our audits provide a reasonable basis for our opinion.

Critical Audit Matters

The critical audit matters communicated below are matters arising from the current period audit of the financial statements that were communicated or required to be communicated to the audit committee and that: (1) relate to accounts or disclosures that are material to the financial statements and (2) involved our especially challenging, subjective or complex judgments. The communication of critical audit matters does not alter in any way our opinion on the consolidated financial statements, taken as a whole, and we are not, by communicating the critical audit matters below, providing separate opinions on the critical audit matters or on the accounts or disclosures to which they relate.

<i>Description of the Matter</i>	Sales deductions As of December 31, 2025, the Company recorded accrued sales deductions of \$10.6 billion. As described in Note 1 to the financial statements under the caption “Product sales and sales deductions,” revenues from product sales are recognized net of accruals for estimated rebates, wholesaler chargebacks, discounts and other deductions (collectively sales deductions), which are established at the time of sale. Auditing the estimation of sales deductions, specifically estimated chargebacks, commercial rebates, and Medicaid rebates related to U.S. product sales, which are netted against product sales, is complex, requires significant judgment, and the amounts involved are material to the financial statements taken as a whole. Revenue from product sales is recognized upon transfer of control of a product to a customer, generally upon delivery, and is based on an amount that reflects the consideration to which the Company expects to be entitled, which represents an amount that is net of accruals for estimated sales deductions. The estimated sales deductions are based on current contractual and statutory requirements, market events and trends, internal and external historical data, and forecasted customer buying patterns.
<i>How We Addressed the Matter in Our Audit</i>	We obtained an understanding, evaluated the design and tested the operating effectiveness of internal controls over the sales deduction processes. This included testing controls over management’s review of significant assumptions and inputs used in the estimate of sales deductions, including actual sales, contractual terms, historical experience, wholesaler inventory levels, demand data and estimated patient population. We also tested management’s controls over the accuracy of forecasting demand activity as well as the completeness and accuracy of the significant components included in the final sales deduction estimates. To test management’s estimated sales deductions, we obtained management’s calculations for the respective estimates and performed the following procedures, among others. We tested management’s estimation process over the determination of sales discount accruals by developing an independent expectation of the estimated accrual balances, including comparing accrual balances recorded by management to those implied by historical payment trends, evaluating trends in actual sales and discount accrual balances, testing a sample of credits issued and payments made throughout the year, and agreeing rates to underlying contract terms.

Description of the Matter

Unrecognized tax benefits

As discussed in Notes 1 and 7 to the consolidated financial statements, the Company operates in various jurisdictions in which differing interpretations of complex tax laws and regulations create uncertainty and necessitate the use of significant judgment in the determination of the Company's unrecognized tax benefits, particularly in the U.S. federal tax jurisdiction where the Company has significant assets and operations. In this regard, the Company uses significant judgment in (1) determining whether a tax position's technical merits are more-likely-than-not to be sustained and (2) measuring the amount of tax benefit that qualifies for recognition. As of December 31, 2025, the Company accrued \$4.4 billion of gross unrecognized tax benefits. Auditing the assessment of the technical merits and measurement of the Company's unrecognized tax benefits is challenging due to the high degree of estimation and management judgment, given the ultimate resolution is dependent on uncontrollable factors such as the resolution of audit disputes with the IRS and results of the U.S. Tax Court case.

How We Addressed the Matter in Our Audit

We obtained an understanding, evaluated the design and tested the operating effectiveness of internal controls over the Company's process to assess the technical merits of its tax positions, as well as management's process to measure the unrecognized tax benefits of those tax positions, particularly in regard to matters in dispute with the IRS. This included testing controls over management's review of the inputs, calculations, assumptions and methods selected to measure the amount of tax benefits that qualify for recognition.

We involved tax controversy and transfer pricing subject matter professionals to assist in assessing the technical merits and measurement of certain of the Company's unrecognized tax benefits. Depending on the nature of the specific tax position and, as applicable, developments with the relevant tax authorities, our procedures included obtaining and reviewing the Company's correspondence with such tax authorities and evaluating certain third-party advice to support the Company's evaluations and recorded positions. We evaluated the status of the ongoing U.S. Tax Court case and developments in the applicable regulatory environments to assess potential effects on the Company's recorded positions. We assessed management's consideration of current tax controversy, litigation and tax litigation trends. We analyzed the assumptions and data used by the Company when it determined the amount of tax benefits to recognize, including applicable interest and penalties, and we tested the accuracy of those underlying calculations. We have also evaluated the Company's income tax disclosures included in Note 7 in relation to these matters.

/s/ Ernst & Young LLP

We have served as the Company's auditor since 1980.
Los Angeles, California
February 13, 2026

AMGEN INC.
CONSOLIDATED STATEMENTS OF INCOME
Years ended December 31, 2025, 2024 and 2023
(In millions, except per-share data)

	2025	2024	2023
Revenues:			
Product sales	\$ 35,148	\$ 32,026	\$ 26,910
Other revenues	1,603	1,398	1,280
Total revenues	<u>36,751</u>	<u>33,424</u>	<u>28,190</u>
Operating expenses:			
Cost of sales	12,037	12,858	8,451
Research and development	7,272	5,964	4,784
Selling, general and administrative	7,050	7,096	6,179
Other	1,312	248	879
Total operating expenses	<u>27,671</u>	<u>26,166</u>	<u>20,293</u>
Operating income	9,080	7,258	7,897
Other income (expense):			
Interest expense, net	(2,755)	(3,155)	(2,875)
Other income, net	<u>2,651</u>	<u>506</u>	<u>2,833</u>
Income before income taxes	8,976	4,609	7,855
Provision for income taxes	<u>1,265</u>	<u>519</u>	<u>1,138</u>
Net income	<u>\$ 7,711</u>	<u>\$ 4,090</u>	<u>\$ 6,717</u>
Earnings per share:			
Basic	\$ 14.33	\$ 7.62	\$ 12.56
Diluted	\$ 14.23	\$ 7.56	\$ 12.49
Weighted-average shares used in the calculation of earnings per share:			
Basic	538	537	535
Diluted	542	541	538

See accompanying notes.

AMGEN INC.
CONSOLIDATED STATEMENTS OF COMPREHENSIVE INCOME
Years ended December 31, 2025, 2024 and 2023
(In millions)

	2025	2024	2023
Net income	\$ 7,711	\$ 4,090	\$ 6,717
Other comprehensive (loss) income, net of reclassification adjustments and taxes:			
Gains (losses) on foreign currency translation adjustments	172	(76)	50
(Losses) gains on cash flow hedges	(377)	309	(150)
Other	13	(10)	42
Other comprehensive (loss) income, net of reclassification adjustments and taxes	(192)	223	(58)
Comprehensive income	<u>\$ 7,519</u>	<u>\$ 4,313</u>	<u>\$ 6,659</u>

See accompanying notes.

AMGEN INC.
CONSOLIDATED BALANCE SHEETS
December 31, 2025 and 2024
(In millions, except per-share data)

	2025	2024
ASSETS		
Current assets:		
Cash and cash equivalents	\$ 9,129	\$ 11,973
Trade receivables, net	9,570	6,782
Inventories	6,225	6,998
Other current assets	4,133	3,277
Total current assets	29,057	29,030
Property, plant and equipment, net	7,913	6,543
Intangible assets, net	22,276	27,699
Goodwill	18,680	18,637
Other noncurrent assets	12,660	9,930
Total assets	<u>\$ 90,586</u>	<u>\$ 91,839</u>
LIABILITIES AND STOCKHOLDERS' EQUITY		
Current liabilities:		
Accounts payable	\$ 2,367	\$ 1,908
Accrued liabilities	18,523	17,641
Current portion of long-term debt	4,599	3,550
Total current liabilities	25,489	23,099
Long-term debt	50,005	56,549
Long-term deferred tax liabilities	1,366	1,616
Long-term tax liabilities	2,690	2,349
Other noncurrent liabilities	2,378	2,349
Contingencies and commitments (see Note 20)		
Stockholders' equity:		
Common stock and additional paid-in capital; \$0.0001 par value per share; 2,750.0 shares authorized; outstanding—538.8 shares in 2025 and 536.9 shares in 2024	34,023	33,533
Accumulated deficit	(25,107)	(27,590)
Accumulated other comprehensive loss	(258)	(66)
Total stockholders' equity	8,658	5,877
Total liabilities and stockholders' equity	<u>\$ 90,586</u>	<u>\$ 91,839</u>

See accompanying notes.

AMGEN INC.

CONSOLIDATED STATEMENTS OF STOCKHOLDERS' EQUITY

Years ended December 31, 2025, 2024 and 2023

(In millions, except per-share data)

	Number of shares of common stock	Common stock and additional paid-in capital	Accumulated deficit	Accumulated other comprehensive loss	Total
Balance as of December 31, 2022	534.0	\$ 32,514	\$ (28,622)	\$ (231)	\$ 3,661
Net income	—	—	6,717	—	6,717
Other comprehensive loss, net of taxes	—	—	—	(58)	(58)
Dividends declared on common stock (\$8.64 per share)	—	—	(4,644)	—	(4,644)
Issuance of common stock in connection with equity award programs	1.4	95	—	—	95
Stock-based compensation expense	—	454	—	—	454
Equity awards issued for Horizon acquisition, net	—	141	—	—	141
Tax impact related to employee stock-based compensation expense	—	(134)	—	—	(134)
Balance as of December 31, 2023	535.4	33,070	(26,549)	(289)	6,232
Net income	—	—	4,090	—	4,090
Other comprehensive income, net of taxes	—	—	—	223	223
Dividends declared on common stock (\$9.13 per share)	—	—	(4,931)	—	(4,931)
Issuance of common stock in connection with equity award programs	2.2	189	—	—	189
Stock-based compensation expense	—	530	—	—	530
Tax impact related to employee stock-based compensation expense	—	(256)	—	—	(256)
Repurchases of common stock	(0.7)	—	(200)	—	(200)
Balance as of December 31, 2024	536.9	33,533	(27,590)	(66)	5,877
Net income	—	—	7,711	—	7,711
Other comprehensive loss, net of taxes	—	—	—	(192)	(192)
Dividends declared on common stock (\$9.66 per share)	—	—	(5,228)	—	(5,228)
Issuance of common stock in connection with equity award programs	1.9	191	—	—	191
Stock-based compensation expense	—	494	—	—	494
Tax impact related to employee stock-based compensation expense	—	(195)	—	—	(195)
Balance as of December 31, 2025	538.8	\$ 34,023	\$ (25,107)	\$ (258)	\$ 8,658

See accompanying notes.

AMGEN INC.
CONSOLIDATED STATEMENTS OF CASH FLOWS
Years ended December 31, 2025, 2024 and 2023
(In millions)

	2025	2024	2023
Cash flows from operating activities:			
Net income	\$ 7,711	\$ 4,090	\$ 6,717
Noncash adjustments to reconcile net income to net cash provided by operating activities:			
Depreciation, amortization and other	5,167	5,592	4,071
Impairment of intangible assets	1,200	159	851
Stock-based compensation expense	494	530	431
Deferred income taxes	(721)	(1,228)	(1,273)
(Gains) losses on equity securities	(2,064)	159	(1,565)
Other items, net	149	(177)	(277)
Changes in operating assets and liabilities, net of acquisitions:			
Trade receivables, net	(2,676)	441	(1,015)
Inventories	886	2,532	491
Other assets	(1,267)	(652)	(564)
Accounts payable	428	312	(402)
Accrued income taxes, net	(2,068)	(1,011)	(1,031)
Long-term tax liabilities	311	(492)	371
Accrued liabilities	499	92	953
Accrued sales incentives and allowance	2,065	1,194	935
Other liabilities	(156)	(51)	(222)
Net cash provided by operating activities	9,958	11,490	8,471
Cash flows from investing activities:			
Cash paid for acquisitions, net of cash acquired	(53)	—	(26,989)
Proceeds from sales of marketable securities	—	—	1,123
Proceeds from maturities of marketable securities	—	—	550
Purchases of property, plant and equipment	(1,858)	(1,096)	(1,112)
Other	(32)	50	224
Net cash used in investing activities	(1,943)	(1,046)	(26,204)
Cash flows from financing activities:			
Net proceeds from issuance of debt	—	—	27,777
Extinguishment of debt	(683)	(659)	(647)
Repayment of debt	(5,000)	(3,600)	(1,454)
Repurchases of common stock	—	(200)	—
Dividends paid	(5,124)	(4,832)	(4,556)
Other	(52)	(124)	(72)
Net cash (used in) provided by financing activities	(10,859)	(9,415)	21,048
(Decrease) increase in cash and cash equivalents	(2,844)	1,029	3,315
Cash and cash equivalents at beginning of year	11,973	10,944	7,629
Cash and cash equivalents at end of year	\$ 9,129	\$ 11,973	\$ 10,944

See accompanying notes.

AMGEN INC.

NOTES TO CONSOLIDATED FINANCIAL STATEMENTS

December 31, 2025

1. Summary of significant accounting policies

Business

Amgen Inc. (including its consolidated subsidiaries, referred to as “Amgen,” “the Company,” “we,” “our” or “us”) is a global biotechnology pioneer that discovers, develops, manufactures and delivers innovative human therapeutics. We operate our business in one operating segment: human therapeutics. See Note 2, Segment and other information.

Principles of consolidation

The consolidated financial statements include the accounts of Amgen and its majority-owned subsidiaries. In determining whether we are the primary beneficiary of a variable interest entity, we consider whether we have both the power to direct activities of the entity that most significantly impact the entity’s economic performance and the obligation to absorb losses of, or the right to receive benefits from, the entity that could potentially be significant to that entity. We do not have any significant interests in any variable interest entities of which we are the primary beneficiary. All material intercompany transactions and balances have been eliminated in consolidation. Certain reclassifications have been made to prior periods in the consolidated financial statements and accompanying notes to conform with the current presentation.

Use of estimates

The preparation of consolidated financial statements in conformity with GAAP requires management to make estimates and assumptions that affect the amounts reported in the consolidated financial statements and accompanying notes. Actual results may differ from those estimates.

Revenues

Product sales and sales deductions

Revenue from product sales is recognized upon transfer of control of a product to a customer, generally upon delivery, based on an amount that reflects the consideration to which we expect to be entitled, net of accruals for estimated rebates, wholesaler chargebacks, discounts and other deductions (collectively, sales deductions) and returns established at the time of sale.

We analyze the adequacy of our accruals for sales deductions quarterly. Amounts accrued for sales deductions are adjusted when trends or significant events indicate that an adjustment is appropriate. Accruals are also adjusted to reflect actual results. Accruals for sales deductions are based primarily on estimates of the amounts earned or to be claimed on the related sales. These estimates take into consideration current contractual and statutory requirements, specific known market events and trends, internal and external historical data and forecasted customer buying patterns. Sales deductions are substantially product specific and therefore, for any given period, can be affected by the mix of products sold. Included in sales deductions are immaterial net adjustments related to prior-period sales due to changes in estimates.

Returns are estimated through comparison of historical return data with their related sales on a production lot basis. Historical rates of return are determined for each product and are adjusted for known or expected changes in the marketplace specific to each product, when appropriate. Historically, sales return provisions have amounted to less than 1% of gross product sales. Changes in estimates for prior-period sales return provisions have historically been immaterial.

Our payment terms vary by types and locations of customers and by products or services offered. Payment terms differ by jurisdiction and customer, but payment is generally required in a term ranging from 30 to 120 days from date of shipment or satisfaction of the performance obligation. For certain products or services and certain customer types, we may require payment before products are delivered or services are rendered to customers.

Indirect taxes collected from customers and remitted to government authorities that are related to sales of the Company’s products, primarily in Europe, are excluded from revenues.

As a practical expedient, sales commissions are expensed when incurred because the amortization period would have been one year or less. These costs are recorded in SG&A expense in the Consolidated Statements of Income.

Other revenues

Other revenues consist primarily of royalty income and corporate partner revenues. Royalties from licensees are based on third-party sales of licensed products and are recorded when the related third-party product sale occurs. Royalty income is estimated based on historical and forecasted sales trends. Corporate partner revenues are composed mainly of license fees and milestones earned and our share of commercial profits generated from collaborations. See Arrangements with multiple-performance obligations, discussed below.

Arrangements with multiple-performance obligations

From time to time, we enter into arrangements for the R&D, manufacture and/or commercialization of products and product candidates. Such arrangements may require us to deliver various rights, services and/or goods, including intellectual property rights/licenses, R&D services, manufacturing services and/or commercialization services. The underlying terms of these arrangements generally provide for consideration to Amgen in the form of nonrefundable, upfront license fees; development and commercial-performance milestone payments; royalty payments; and/or profit sharing.

In arrangements involving more than one performance obligation, each required performance obligation is evaluated to determine whether it qualifies as a distinct performance obligation based on whether (i) the customer can benefit from the good or service either on its own or together with other resources that are readily available and (ii) the good or service is separately identifiable from other promises in the contract. The consideration under the arrangement is then allocated to each separate distinct performance obligation based on its respective relative stand-alone selling price. The estimated selling price of each deliverable reflects our best estimate of what the selling price would be if the deliverable was regularly sold by us on a stand-alone basis or by using an adjusted market assessment approach if selling price on a stand-alone basis is not available.

The consideration allocated to each distinct performance obligation is recognized as revenue when control of the related goods or services is transferred. Consideration associated with at-risk substantive performance milestones is recognized as revenue when it is probable that a significant reversal of the cumulative revenue recognized will not occur. We utilize the sales- and usage-based royalty exception in arrangements that resulted from the license of intellectual property, recognizing revenues generated from royalties or profit sharing as the underlying sales occur.

Research and development costs

R&D costs are expensed as incurred and primarily include salaries, benefits and other staff-related costs; facilities and overhead costs; clinical trial and related clinical manufacturing costs; contract services and other outside costs; information systems' costs; and amortization of acquired technology used in R&D with alternative future uses. R&D expenses also include costs and cost recoveries associated with third-party R&D arrangements, including upfront fees and milestones paid to third parties in connection with technologies that had not reached technological feasibility and did not have an alternative future use. Net payment or reimbursement of R&D costs is recognized when the obligations are incurred or as we become entitled to the cost recovery. See Note 9, Collaborations.

Selling, general and administrative costs

SG&A costs are primarily composed of salaries, benefits and other staff-related costs associated with sales and marketing, finance, legal and other administrative personnel; facilities and overhead costs; outside marketing, advertising and legal expenses; the U.S. healthcare reform federal excise fee on Branded Prescription Pharmaceutical Manufacturers and Importers; and other general and administrative costs. Advertising costs are expensed as incurred and were \$963 million, \$987 million and \$647 million during the years ended December 31, 2025, 2024 and 2023, respectively. SG&A expenses also include costs and cost recoveries associated with marketing and promotion efforts under certain collaborative arrangements. Net payment or reimbursement of SG&A costs is recognized when the obligations are incurred or we become entitled to the cost recovery. See Note 9, Collaborations.

Leases

At inception of a contract, we determine whether an arrangement is or contains a lease. For all leases, we determine the classification as either operating or financing. Operating leases are included in Other noncurrent assets, Accrued liabilities and Other noncurrent liabilities in our Consolidated Balance Sheets.

ROU assets represent our right to use an underlying asset for the lease term, and lease liabilities represent our obligation to make lease payments under the lease. Lease recognition occurs at the commencement date, and lease liability amounts are based on the present value of lease payments made during the lease term. Our lease terms may include options to extend or terminate a lease when it is reasonably certain that we will exercise that option. Because most of our leases do not provide information to determine an implicit interest rate, we use our incremental borrowing rate in determining the present value of

lease payments. ROU assets also include any lease payments made prior to the commencement date less lease incentives received. Operating lease expense is recognized on a straight-line basis over the lease term.

We have lease agreements with both lease and nonlease components, which are generally accounted for together as a single lease component. In addition, for certain vehicle and equipment leases, we apply a portfolio approach to determine the lease term and discount rate.

Stock-based compensation

We have stock-based compensation plans under which various types of equity-based awards are granted, including RSUs, performance units and stock options. The fair values of RSUs and stock option awards, which are subject only to service conditions with graded vesting, are recognized as compensation expense, generally on a straight-line basis over the service period, net of estimated forfeitures. The fair values of performance unit awards are recognized as compensation expense, generally on a straight-line basis from the grant date to the end of the performance period. See Note 5, Stock-based compensation.

Income taxes

We provide for income taxes based on pretax income and applicable tax rates in the various jurisdictions in which we operate. Significant judgment is required in determining our provision for income taxes and income tax assets and liabilities, including evaluating uncertainties in the application of accounting principles and complex tax laws. Deferred income taxes are recorded for the expected tax consequences of temporary differences between the bases of assets and liabilities, as well as for loss and tax credit carryforwards for financial reporting purposes and amounts recognized for income tax purposes. We record a valuation allowance to reduce our deferred tax assets to the amount of future tax benefit that is more likely than not to be realized.

We recognize the tax benefit from an uncertain tax position only if it is more likely than not that the tax position will be sustained upon examination by tax authorities based on the technical merits of the position. The tax benefit recognized in the consolidated financial statements for a particular tax position is based on the largest benefit that is more likely than not to be realized. The amount of UTBs is adjusted as appropriate for changes in facts and circumstances, such as significant amendments to existing tax law, new regulations or interpretations by tax authorities, new information obtained during a tax examination or resolution of an examination. We recognize both accrued interest and penalties, when appropriate, related to UTBs in income tax expense. See Note 7, Income taxes.

Amgen is subject to current U.S. tax on the earnings of our foreign subsidiaries. We previously established deferred taxes related to this U.S. tax, which requires us to recognize deferred taxes for temporary basis differences expected to reverse and be subject to this tax in future years. These are ongoing adjustments that are likely to occur in future periods.

Acquisitions

We first determine whether a set of assets acquired constitutes a business and should be accounted for as a business combination. If the assets acquired do not constitute a business, we account for the transaction as an asset acquisition. Business combinations are accounted for by means of the acquisition method of accounting. Under the acquisition method, assets acquired, including IPR&D projects, and liabilities assumed are recorded in our consolidated financial statements at their respective fair values as of the acquisition date. The excess of the fair value of consideration transferred over the fair value of the net assets acquired is recorded as goodwill. Contingent consideration obligations incurred in connection with a business combination, including the assumption of an acquiree's liability arising from an acquisition it consummated prior to our acquisition, are recorded at their fair values on the acquisition date and remeasured at their fair values each subsequent reporting period until the related contingencies have been resolved. The resulting changes in fair values are recorded in earnings. In contrast, asset acquisitions are accounted for by using a cost accumulation and allocation model. Under this model, the cost of the acquisition is allocated to the assets acquired and liabilities assumed. IPR&D projects with no alternative future use are recorded in R&D expense upon acquisition, and contingent consideration obligations incurred in connection with an asset acquisition are recorded when it is probable that they will occur and they can be reasonably estimated. See Note 4, Acquisition, and Note 18, Fair value measurement.

Cash equivalents

We consider cash equivalents to be only those investments that are highly liquid, that are readily convertible to cash and that mature within three months from the date of purchase.

Interest-bearing securities

We consider our interest-bearing securities investment portfolio as available-for-sale, and accordingly, these investments are recorded at fair value, with unrealized gains and losses recorded in AOCI. Investments with maturities beyond one year may be classified as short-term marketable securities in the Consolidated Balance Sheets due to their highly liquid nature and because they represent the Company's investments that are available for current operations. See Note 10, Investments, and Note 18, Fair value measurement.

Inventories

Inventories are stated at the lower of cost or net realizable value. Cost, which includes amounts related to materials, labor and overhead, is determined in a manner that approximates the first-in, first-out method. Net realizable value is the estimated selling price in the ordinary course of business less reasonably predictable costs of completion, disposal and transportation. See Note 11, Inventories.

Derivatives

We recognize all of our derivative instruments as either assets or liabilities at fair value in the Consolidated Balance Sheets. The accounting for changes in the fair value of a derivative instrument depends on whether the derivative has been formally designated and qualifies as part of a hedging relationship under the applicable accounting standards and, further, on the type of hedging relationship. For derivatives formally designated as hedges, we assess both at inception and quarterly thereafter whether the hedging derivatives are highly effective in offsetting changes in either the fair value or cash flows of the hedged item. Our derivatives that are not designated and do not qualify as hedges are adjusted to fair value through current earnings. See Note 18, Fair value measurement, and Note 19, Derivative instruments.

Property, plant and equipment, net

Property, plant and equipment is recorded at historical cost, net of accumulated depreciation, amortization and, if applicable, impairment charges. We review our property, plant and equipment assets for impairment whenever events or changes in circumstances indicate that the carrying amount of an asset may not be recoverable. Depreciation is recorded over the assets' useful lives on a straight-line basis. Leasehold improvements are amortized on a straight-line basis over the shorter of their estimated useful lives or lease terms. See Note 12, Property, plant and equipment.

Cloud computing arrangements

We capitalize qualifying implementation costs under cloud computing arrangements (CCA). Capitalized CCA implementation costs are allocated between Other current assets and Other noncurrent assets in the Consolidated Balance Sheets. As of December 31, 2025 and 2024, capitalized CCA implementation costs totaled \$272 million and \$227 million, respectively. For critical software-as-a-service (SaaS) platforms, amortization is recorded over an established four-year period, and for all other arrangements, amortization is recorded over the expected term of the related contract.

Goodwill and other intangible assets

Finite-lived intangible assets are recorded at cost, net of accumulated amortization and, if applicable, impairment charges. Amortization of finite-lived intangible assets is recorded over the assets' estimated useful lives on a straight-line basis or based on the pattern in which economic benefits are consumed, if reliably determinable. We review our finite-lived intangible assets for impairment whenever events or changes in circumstances indicate that the carrying amount of an asset may not be recoverable. See Note 13, Goodwill and other intangible assets.

The fair values of IPR&D projects acquired in a business combination that are not complete are capitalized and accounted for as indefinite-lived intangible assets until completion or abandonment of the related R&D efforts. Upon successful completion of the project, the capitalized amount is amortized over its estimated useful life. If a project is abandoned, all remaining capitalized amounts are written off immediately. Major risks and uncertainties are often associated with IPR&D projects because we are required to obtain regulatory approvals before marketing the resulting products. Such approvals require completing clinical trials that demonstrate a product candidate is safe and effective. Consequently, the eventual realized value of the acquired IPR&D project may vary from its fair value at the date of acquisition, and IPR&D impairment charges may occur in future periods.

Capitalized IPR&D projects are reviewed for impairment annually and whenever events or changes in circumstances indicate that the carrying amount may not be recoverable. We consider various factors for potential impairment, including the current legal and regulatory environment and the competitive landscape. Adverse clinical trial results, significant delays in obtaining marketing approval, the inability to bring a product to market and the introduction or advancement of competitors' products could result in partial or full impairment of the related intangible assets.

We perform an impairment test of goodwill annually and whenever events or changes in circumstances indicate that the carrying amount may not be recoverable. To date, an impairment of goodwill has not been recorded. See Note 13, Goodwill and other intangible assets.

Contingencies

In the ordinary course of business, we are involved in various legal proceedings, government investigations and other matters that are complex in nature and have outcomes that are difficult to predict. Certain of these proceedings are discussed in Note 20, Contingencies and commitments. We record accruals for loss contingencies to the extent that we conclude it is probable that a liability has been incurred and the amount of the related loss can be reasonably estimated. We evaluate, on a quarterly basis, developments in legal proceedings and other matters that could cause an increase or decrease in the amount of the liability that has been accrued previously.

Foreign currency translation

The net assets of international subsidiaries whose functional currencies are not in U.S. dollars are translated into U.S. dollars using current exchange rates. The U.S. dollar effects that arise from translation of the net assets of these subsidiaries at changing rates are recognized in AOCI. The subsidiaries' earnings are translated into U.S. dollars by using average exchange rates.

Equity investments

Marketable and nonmarketable equity securities

Investments in publicly traded equity securities with readily determinable fair values are recorded at quoted market prices for identical securities, with changes in fair value recorded in Other income, net, in the Consolidated Statements of Income. Investments in equity securities without readily determinable fair values are recorded at cost less impairment, if any, adjusted for changes resulting from observable price changes in orderly transactions for identical or similar securities. Such adjustments are recorded in Other income, net, in the Consolidated Statements of Income.

Equity method investments

Equity investments that give us the ability to exert significant influence, but not control, over an investee are accounted for under the equity method of accounting, unless we have elected the fair value option. For equity method investments for which the fair value option has been elected, changes in fair value are recorded in Other income, net, in the Consolidated Statements of Income.

Additionally, we hold investments in limited partnerships, which primarily invest in early-stage biotechnology companies. As a practical expedient, such limited partnership investments are measured by using our proportionate share of the net asset values of the underlying investments held by the limited partnerships, with such changes included in Other income, net, in the Consolidated Statements of Income.

Recently adopted accounting pronouncements

In December 2023, the FASB issued ASU No. 2023-09, Income Taxes (Topic 740): Improvements to Income Tax Disclosures, to improve income tax disclosure requirements by requiring more detailed information in several income tax disclosures, such as enhancing disclosure of income taxes paid and requiring disaggregation of the effective income tax rate reconciliation. The standard is effective for public business entities such as Amgen for annual periods beginning after December 15, 2024. Early adoption is permitted, and entities may apply the standard prospectively or retrospectively. We prospectively adopted this standard in fiscal year 2025, which resulted in incremental income taxes disclosures. See Note 7, Income taxes.

Recent accounting pronouncements not yet adopted

In November 2024, the FASB issued ASU No. 2024-03, Income Statement—Reporting Comprehensive Income—Expense Disaggregation Disclosures (Subtopic 220-40): Disaggregation of Income Statement Expenses, to improve disclosures about a public business entity's expenses by requiring disaggregated disclosures of certain types of expenses, including purchases of inventory, employee compensation, depreciation, intangible amortization and depletion, as applicable, for each income statement caption that includes those expenses. In addition, the standard will require entities to define and disclose total selling expenses. The standard is effective for public business entities such as Amgen for annual periods beginning after December 15, 2026, and interim periods beginning after December 15, 2027. Early adoption is permitted, and entities may apply the standard prospectively or retrospectively. We are currently evaluating the impact of adopting this standard on our consolidated financial statements and related disclosures.

In September 2025, the FASB issued ASU No. 2025-06, Intangibles—Goodwill and Other—Internal-Use Software (Subtopic 350-40): Targeted Improvements to the Accounting for Internal-Use Software, to modernize the accounting for software costs, including updating guidance on the recognition and measurement of costs incurred in connection with development and implementation activities related to internal-use software. The standard is effective for all entities for annual periods beginning after December 15, 2027, and interim periods within those annual periods. Early adoption is permitted, and entities may apply the standard prospectively or retrospectively. We evaluated the impact of adopting this new standard on our consolidated financial statements and related disclosures and do not expect the adoption to have a material impact.

2. Segment and other information

We operate our business in one operating segment, which also represents one reportable segment: human therapeutics. Therefore, results of our operations are reported on a consolidated basis for purposes of segment reporting, consistent with internal management reporting.

The human therapeutics segment is engaged in the discovery, development, manufacturing and delivery of innovative medicines to fight some of the world's toughest diseases. The Company's Chief Executive Officer has been identified as the chief operating decision maker (CODM). The CODM manages and allocates resources on a consolidated basis. The determination of a single segment is consistent with the financial information regularly reviewed by the CODM for purposes of evaluating performance and allocating resources, which is reviewed on a consolidated basis.

As the Company's CODM evaluates the financial performance of the Company's human therapeutics segment on a consolidated basis, the measure of segment performance is net income, as reflected in the Consolidated Statements of Income. The CODM uses net income to allocate resources on a consolidated basis, which enables the CODM to assess both the overall level of resources available and optimize distribution of resources across functions, therapeutic areas, regions and R&D programs in line with our long-term corporate-wide strategic goals. In addition, the CODM may also evaluate financial performance based on net income adjusted for certain items that are unusual and non-recurring. As the Company manages its assets on a consolidated basis, the measure of segment assets is total assets, as reflected in the Consolidated Balance Sheets. See Note 10, Investments, for further information regarding equity method investments, and Net cash used in investing activities in the Consolidated Statements of Cash Flows for further information regarding capital expenditures.

The following table provides segment revenues, significant segment expenses, other segment items and reported segment net income for the Company's one reportable segment, as well as a reconciliation of segment net income to the Company's total consolidated net income for the years ended December 31, 2025, 2024 and 2023 (in millions):

	Years ended December 31,		
	2025	2024	2023
Revenues:			
Product sales	\$ 35,148	\$ 32,026	\$ 26,910
Other revenues	1,603	1,398	1,280
Total revenues	36,751	33,424	28,190
Less:			
Manufacturing cost of sales ⁽¹⁾⁽²⁾	9,845	11,118	7,347
Profit share and royalties in cost of sales ⁽¹⁾	2,192	1,740	1,104
Research and development ⁽¹⁾	7,272	5,964	4,784
Sales and marketing ⁽¹⁾	4,590	4,713	3,784
General and administrative ⁽¹⁾	2,460	2,383	2,395
Other segment items ⁽³⁾	(931)	252	(729)
Interest income	(408)	(510)	(1,225)
Interest expense, net	2,755	3,155	2,875
Provision for income taxes	1,265	519	1,138
Segment net income	7,711	4,090	6,717
Reconciliation of profit or loss:			
Adjustments and reconciling items	—	—	—
Consolidated net income	\$ 7,711	\$ 4,090	\$ 6,717

⁽¹⁾ During the years ended December 31, 2025, 2024 and 2023, we recognized amortization expense on our intangible assets of \$4.3 billion, \$4.8 billion and \$3.2 billion, respectively. Amortization of intangible assets is primarily included in Cost of sales in the Consolidated Statements of Income. In addition, during the years ended December 31, 2025, 2024 and 2023, we recognized depreciation and ROU asset amortization expense of \$887 million, \$805 million and \$824 million, respectively.

⁽²⁾ During the years ended December 31, 2025, 2024 and 2023, manufacturing cost of sales included amortization of step-up to fair value of inventory acquired in business combinations of \$1.3 billion, \$2.4 billion and \$656 million, respectively.

⁽³⁾ Other segment items included in Segment net income primarily consists of: (i) fair value adjustments on equity securities (see Note 10, Investments); (ii) net impairment charges on intangible assets (see Note 13, Goodwill and other intangible assets); and (iii) expenses related to restructuring and cost-savings initiatives.

3. Revenues

We operate our business in one operating segment, which also represents one reportable segment: human therapeutics. Therefore, results of our operations are reported on a consolidated basis for purposes of segment reporting, consistent with internal management reporting. Revenues by product and by geographic area, based on customers' locations, are presented below. A substantial portion of ROW product sales relates to products sold in Europe.

Revenues were as follows (in millions):

	Year ended December 31, 2025			Year ended December 31, 2024			Year ended December 31, 2023		
	U.S.	ROW	Total	U.S.	ROW	Total	U.S.	ROW	Total
Prolia	\$ 2,978	\$ 1,436	\$ 4,414	\$ 2,885	\$ 1,489	\$ 4,374	\$ 2,733	\$ 1,315	\$ 4,048
Repatha	1,663	1,353	3,016	1,139	1,083	2,222	793	842	1,635
Otezla	1,839	426	2,265	1,699	427	2,126	1,777	411	2,188
ENBREL	2,199	27	2,226	3,288	28	3,316	3,650	47	3,697
EVENITY	1,600	500	2,100	1,131	432	1,563	809	351	1,160
XGEVA	1,355	729	2,084	1,507	718	2,225	1,527	585	2,112
TEPEZZA ⁽¹⁾	1,758	145	1,903	1,835	16	1,851	441	7	448
BLINCYTO	1,049	510	1,559	800	416	1,216	566	295	861
Nplate	1,027	497	1,524	970	486	1,456	996	481	1,477
TEZSPIRE ⁽²⁾	1,478	—	1,478	972	—	972	567	—	567
KYPROLIS	913	499	1,412	948	555	1,503	921	482	1,403
Aranesp	416	973	1,389	386	956	1,342	452	910	1,362
KRYSTEXXA ⁽¹⁾	1,340	—	1,340	1,185	—	1,185	272	—	272
Vectibix	604	571	1,175	519	526	1,045	461	523	984
Other products ⁽³⁾	5,437	1,826	7,263	4,037	1,593	5,630	3,307	1,389	4,696
Total product sales ⁽⁴⁾	25,656	9,492	35,148	23,301	8,725	32,026	19,272	7,638	26,910
Other revenues	763	840	1,603	562	836	1,398	534	746	1,280
Total revenues	\$ 26,419	\$ 10,332	\$ 36,751	\$ 23,863	\$ 9,561	\$ 33,424	\$ 19,806	\$ 8,384	\$ 28,190

⁽¹⁾ TEPEZZA and KRYSTEXXA were acquired from the acquisition of Horizon on October 6, 2023, and include product sales in the periods after the acquisition date.

⁽²⁾ TEZSPIRE is marketed by our collaborator AstraZeneca outside the United States.

⁽³⁾ Consists of product sales of our non-principal products.

⁽⁴⁾ Hedging gains and losses, which are included in product sales, were not material for the years ended December 31, 2025, 2024 and 2023.

In the United States, we sell primarily to pharmaceutical wholesale distributors that we use as the principal means of distributing our products to healthcare providers. Outside the United States, we sell principally to healthcare providers and/or pharmaceutical wholesale distributors depending on the distribution practice in each country. We monitor the financial condition of our larger customers and limit our credit exposure by setting credit limits and, in certain circumstances, by requiring letters of credit or obtaining credit insurance.

For each of the years ended December 31, 2025, 2024 and 2023, we had product sales to three customers that individually accounted for more than 10% of total revenues. For the year ended December 31, 2025, on a combined basis, these customers accounted for 77% of total gross revenues as shown in the following table. Certain information with respect to these customers was as follows (dollar amounts in millions):

	Years ended December 31,		
	2025	2024	2023
McKesson Corporation:			
Gross product sales	\$ 26,253	\$ 22,173	\$ 19,035
% of total gross revenues	34 %	33 %	33 %
Cencora, Inc.:			
Gross product sales	\$ 20,986	\$ 18,387	\$ 16,625
% of total gross revenues	27 %	27 %	29 %
Cardinal Health, Inc.:			
Gross product sales	\$ 12,562	\$ 11,278	\$ 9,775
% of total gross revenues	16 %	17 %	17 %

As of December 31, 2025 and 2024, amounts due from these three customers each exceeded 10% of gross trade receivables and accounted for 75% and 70%, respectively, of net trade receivables on a combined basis. As of December 31, 2025 and 2024, 21% and 26%, respectively, of net trade receivables were due from customers located outside the United States, the majority of which were from Europe. Our total allowance for doubtful accounts as of December 31, 2025 and 2024, was not material.

4. Acquisition

Acquisition of Horizon Therapeutics plc

On October 6, 2023, Amgen completed its acquisition of Horizon by acquiring all of the outstanding shares of Horizon for \$116.50 per share in cash, representing a total consideration of approximately \$27.8 billion. Horizon is a global biotechnology company focused on the discovery, development and commercialization of medicines that address critical needs of patients impacted by rare, autoimmune and severe inflammatory diseases. The acquisition, which was accounted for as a business combination, aligns with Amgen's core strategy of delivering innovative medicines that make a significant difference for patients suffering from serious diseases and strengthens Amgen's leading rare disease portfolio by adding first-in-class, early-in-lifecycle medicines, including TEPEZZA for TED, KRYSTEXXA for chronic refractory gout and UPLIZNA for neuromyelitis optica spectrum disorder. Upon its acquisition, Horizon became a wholly owned subsidiary of Amgen, and its operations have been included in our consolidated financial statements commencing on the acquisition date.

During the year ended December 31, 2024, the purchase price allocation of the acquisition was completed and measurement period adjustments were finalized, which included changes to the purchase price allocation that resulted in a net increase of approximately \$25 million to goodwill. The measurement period adjustments resulted primarily from adjustments to acquired assets and liabilities, including deferred tax attributes, based on facts and circumstances that existed as of the acquisition date and did not result from events subsequent to the acquisition date. The adjustments did not have a significant impact on Amgen's results of operations during the year ended December 31, 2024, and would not have had a significant impact on prior-period results if the adjustments had been made as of the acquisition date.

The following table summarizes the final total consideration and allocated acquisition date fair values of assets acquired and liabilities assumed, inclusive of measurement-period adjustments (in millions):

Cash and cash equivalents	\$	681
Inventories		5,014
Property, plant and equipment, net		318
Finite-lived intangible assets—developed-product-technology rights		19,590
IPR&D		1,060
Goodwill		3,136
Deferred tax asset		795
Deferred tax liability		(2,488)
Other assets and liabilities, net		(273)
Total assets acquired, net of liabilities assumed	\$	<u>27,833</u>

The \$27.8 billion total consideration for this transaction consisted of (i) cash consideration transferred to common shareholders of \$26.7 billion; (ii) cash consideration transferred to vested and outstanding options, outstanding RSU awards and outstanding performance share unit awards of \$523 million; (iii) fair value of Amgen replacement awards (based on conversion of outstanding employee RSU awards) of \$180 million representing noncash consideration; and (iv) a portion of Horizon's debt, settled by Amgen on the acquisition closing date, of \$382 million. Amgen issued 1.7 million replacement equity awards with the original vesting conditions, the fair value of which was determined based on the acquisition date fair value based on the conversion calculation. See Note 5, Stock-based compensation.

The estimated fair values of \$20.7 billion for the developed-product-technology rights and IPR&D intangible assets were determined using a multi-period excess earnings income approach that discounts expected future cash flows to present value by applying a discount rate that represents the estimated rate that market participants would use to value the intangible assets. The projected cash flows were based on certain assumptions attributable to the respective intangible asset, including estimates of future revenues and expenses, the time and resources needed to complete development and the probabilities of obtaining marketing approval from the FDA and other regulatory agencies. The developed-product-technology rights are being amortized on a straight-line basis over a weighted-average period of approximately 10 years from the acquisition date using the straight-line methodology.

The estimated fair value of the acquired inventory of \$5.0 billion was determined using the comparative sales method, which uses actual or expected selling prices of inventory as the base amount to which adjustments for selling effort and a profit on the buyer's effort are applied. The inventory fair value adjustment is being amortized using a weighted-average inventory turnover, which we estimate to approximate 27 months from the acquisition date.

A deferred tax liability of \$2.5 billion was recognized on the temporary differences related to the book bases and tax bases of the acquired identifiable assets and assumed liabilities, primarily driven by the intangible assets acquired, as well as associated deferred tax asset for anticipatory foreign tax credits of \$795 million.

The excess of the acquisition date consideration over the fair values assigned to the assets acquired and the liabilities assumed of \$3.1 billion was recorded as goodwill, which is not deductible for tax purposes. The goodwill value represents expected synergies from the marketed products acquired and other benefits.

During the three months ended December 31, 2023, the Company incurred approximately \$487 million of acquisition costs related to the closing of our Horizon acquisition, consisting of \$167 million for share-based payments to settle non-vested equity awards attributable to post-combination services, severance and other employee-related expenses and \$320 million for transaction costs. These costs were included primarily in SG&A expense in the Consolidated Statements of Income.

Supplemental Pro Forma Financial Information

The following table presents the unaudited supplemental pro forma results of a hypothetical combined Amgen and Horizon entity for the years ended December 31, 2023 and 2022, as if the acquisition of Horizon had occurred on January 1, 2022 (in millions):

	Years ended December 31,	
	2023	2022
Total revenues	\$ 30,969	\$ 29,964
Net income	\$ 5,383	\$ 2,381

The unaudited supplemental pro forma combined financial information was prepared using the acquisition method of accounting and was based on the historical financial information of Amgen and Horizon. In order to reflect the occurrence of the acquisition on January 1, 2022, the unaudited supplemental pro forma financial information includes adjustments to reflect: (i) incremental amortization expense based on the fair values of the identifiable intangible assets and inventory step-up; (ii) the additional interest expense associated with the issuance of debt to finance the acquisition; (iii) the reclassification of transaction and other acquisition-related costs incurred during the three months ended December 31, 2023, to the year ended December 31, 2022; and (iv) the income tax impact using an estimated effective tax rate applied to the combined entity. The unaudited supplemental pro forma financial information is not necessarily indicative of what the consolidated results of operations would have been had the acquisition been completed on January 1, 2022. In addition, the unaudited pro forma financial information is not a projection of future results of operations of the combined company, nor does it reflect the expected realization of any synergies or cost savings associated with the acquisition.

5. Stock-based compensation

Our Amended 2009 Plan authorizes for issuance to employees of Amgen and nonemployee members of our Board of Directors shares of our common stock pursuant to grants of equity-based awards, including RSUs, stock options and performance units. The pool of shares available under the Amended 2009 Plan is reduced by one share for each stock option granted and by 1.9 shares for other types of awards granted, including full-value awards. In general, if any shares subject to an award granted under the Amended 2009 Plan expire or become forfeited, terminated or canceled without the issuance of shares, the shares subject to such awards are added back into the authorized pool on the same basis that they were removed. In addition, under the Amended 2009 Plan, shares withheld to pay for minimum statutory tax obligations with respect to full-value awards are added back into the authorized pool on the basis of 1.9 shares. As of December 31, 2025, the Amended 2009 Plan provides for future grants and/or issuances of up to approximately 35 million shares of our common stock. Stock-based awards under our employee compensation plans are made with newly issued shares reserved for this purpose.

The following table reflects the components of stock-based compensation expense recognized in our Consolidated Statements of Income (in millions):

	Years ended December 31,		
	2025	2024	2023
RSUs	\$ 333	\$ 351	\$ 309
Performance units	112	133	121
Stock options	49	46	43
Total stock-based compensation expense, pretax	494	530	473
Tax benefit from stock-based compensation expense	(106)	(114)	(102)
Total stock-based compensation expense, net of tax	\$ 388	\$ 416	\$ 371

Restricted stock units and stock options

Eligible employees generally receive an annual grant of RSUs and, for certain executive-level employees, stock options, with the size and type of award generally determined by the employee's salary grade and performance level. Certain management and professional-level employees typically receive RSU grants upon commencement of employment. Nonemployee members of our Board of Directors also receive an annual grant of RSUs.

Our RSU and stock option grants provide for accelerated or continued vesting in certain circumstances as defined in the plans and related grant agreements, including upon death, disability, termination in connection with a change in control and the retirement of employees who meet certain service and/or age requirements. For awards granted prior to April 1, 2025, RSUs and stock options generally vest in equal amounts on the second, third and fourth anniversaries of the grant date. Effective for awards granted on or after April 1, 2025, RSUs and stock options generally vest in equal amounts on the first, second, third and fourth anniversaries of the grant date. RSUs accrue dividend equivalents, which are typically payable in shares only when and to the extent the underlying RSUs vest and are issued to the recipient.

Restricted stock units

The grant date fair value of an RSU equals the closing price of our common stock on the grant date, as RSUs accrue dividend equivalents during their vesting period, except with respect to certain holders of Horizon unvested RSUs who were granted replacement Amgen RSUs in 2023 under the original terms of the awards in connection with the Horizon acquisition based on the terms of the transaction. See Note 4, Acquisition. Subsequent to the Horizon acquisition, \$42 million of the RSUs were accelerated and cash settled in 2023. The weighted-average grant date fair values per unit of RSUs granted (excluding replacement awards granted to Horizon RSU holders) during the years ended December 31, 2025, 2024 and 2023, were \$273.10, \$301.36 and \$237.70, respectively.

The following table summarizes information regarding our RSUs:

	Year ended December 31, 2025	
	Units (in millions)	Weighted-average grant date fair value
Balance nonvested as of December 31, 2024	3.5	\$ 265.07
Granted	1.6	\$ 273.10
Vested	(1.2)	\$ 243.73
Forfeited	(0.2)	\$ 277.14
Balance nonvested as of December 31, 2025	3.7	\$ 274.23

The total grant date fair values of RSUs that vested during the years ended December 31, 2025, 2024 and 2023, were \$283 million, \$401 million and \$309 million, respectively.

Stock options

The exercise price of stock options is set as the closing price of our common stock on the grant date, and the related number of shares granted is fixed at that point in time. Awards expire 10 years from the date of grant. We use the Black–Scholes option valuation model to estimate the grant date fair value of stock options.

The weighted-average assumptions used in the option valuation model and the resulting weighted-average grant date fair values of stock options granted were as follows:

	Years ended December 31,		
	2025	2024	2023
Closing price of our common stock on grant date	\$ 270.44	\$ 300.30	\$ 235.97
Expected volatility (average of implied and historical volatility)	27.8 %	26.9 %	23.3 %
Expected life (in years)	5.7	5.7	5.7
Risk-free interest rate	4.0 %	4.4 %	3.4 %
Expected dividend yield	3.2 %	3.2 %	3.5 %
Fair value of stock options granted	\$ 62.76	\$ 69.34	\$ 41.86

The following table summarizes information regarding our stock options:

	Year ended December 31, 2025			
	Options (in millions)	Weighted- average exercise price	Weighted- average remaining contractual life (in years)	Aggregate intrinsic value (in millions)
Balance unexercised as of December 31, 2024	5.9	\$ 225.84		
Granted	0.9	\$ 270.44		
Exercised	(0.6)	\$ 202.34		
Expired/forfeited	(0.1)	\$ 258.80		
Balance unexercised as of December 31, 2025	6.1	\$ 234.45	6.0	\$ 563
Vested or expected to vest as of December 31, 2025	5.9	\$ 233.42	5.9	\$ 555
Exercisable as of December 31, 2025	3.5	\$ 211.48	4.4	\$ 401

The total intrinsic values of options exercised during the years ended December 31, 2025, 2024 and 2023, were \$70 million, \$70 million and \$33 million, respectively. The actual tax benefits realized from tax deductions from option exercises during the years ended December 31, 2025, 2024 and 2023, were \$15 million, \$15 million and \$7 million, respectively.

As of December 31, 2025, \$537 million of unrecognized compensation cost was related to nonvested RSUs and unvested stock options, which is expected to be recognized over a weighted-average period of 1.7 years.

Performance units

Certain management-level employees also receive annual grants of performance units, which give the recipient the right to receive common stock that is contingent upon achievement of specified preestablished goals over the performance period, which is generally three years. The performance goals for the units granted during the years ended December 31, 2025, 2024 and 2023, which are accounted for as equity awards, are based on (i) Amgen's total stockholder return compared with a comparator group of companies, which are considered market conditions and are therefore reflected in the grant date fair values of the units, and (ii) Amgen's stand-alone financial performance measures, which are considered performance conditions. The expense recognized for awards is based on the grant date fair value of a unit multiplied by the number of units expected to be earned with respect to the related performance conditions, net of estimated forfeitures. Depending on the outcome of these performance goals, a recipient may ultimately earn a number of units greater or less than the number of units granted. Shares of our common stock are issued on a one-for-one basis for each performance unit earned. In general, performance unit awards vest at the end of the performance period. The performance award program provides for accelerated or continued vesting in certain circumstances as defined in the plan, including upon death, disability, a change in control and retirement of employees who meet certain service and/or age requirements. Performance units accrue dividend equivalents that are typically payable in shares only when and to the extent the underlying performance units vest and are issued to the recipient, including with respect to market and performance conditions that affect the number of performance units earned.

We use a payout simulation model to estimate the grant date fair value of performance units. The weighted-average assumptions used in the payout simulation model and the resulting weighted-average grant date fair values of performance units granted were as follows:

	Years ended December 31,		
	2025	2024	2023
Closing price of our common stock on grant date	\$ 270.44	\$ 300.30	\$ 235.97
Volatility	24.2 %	22.1 %	21.6 %
Risk-free interest rate	3.7 %	4.6 %	3.7 %
Fair value of units granted	\$ 290.24	\$ 321.61	\$ 252.49

The payout simulation model assumes correlations of returns of the stock prices of our common stock and the common stocks of the comparator groups of companies and stock price volatilities of the comparator groups of companies to simulate stockholder returns over the performance periods and their resulting impact on the payout percentages based on the contractual terms of the performance units.

As of December 31, 2025 and 2024, 1.2 million and 1.4 million performance units were outstanding, respectively, with weighted-average grant date fair values per unit of \$278.62 and \$263.86 per unit, respectively. During the year ended December 31, 2025, 0.3 million performance units with a weighted-average grant date fair value per unit of \$290.24 were granted, and 0.1 million performance units with a weighted-average grant date fair value per unit of \$278.87 were forfeited.

The total fair values of performance units paid during the years ended December 31, 2025, 2024 and 2023, were \$163 million, \$182 million and \$109 million, respectively, based on the number of performance units earned multiplied by the closing stock price of our common stock on the last day of the performance period.

As of December 31, 2025, \$64 million of unrecognized compensation cost was related to nonvested performance units, which is expected to be recognized over a weighted-average period of one year.

6. Defined contribution plan

The Company has defined contribution plans to which certain employees of the Company and participating subsidiaries may defer compensation for income tax purposes. Participants are eligible to receive matching contributions based on their contributions, in addition to other Company contributions. Defined contribution plan expenses were \$407 million, \$375 million and \$311 million for the years ended December 31, 2025, 2024 and 2023, respectively.

7. Income taxes

Income before income taxes included the following (in millions):

	Years ended December 31,		
	2025	2024	2023
Domestic	\$ 8,220	\$ 4,040	\$ 4,047
Foreign	756	569	3,808
Total income before income taxes	<u>\$ 8,976</u>	<u>\$ 4,609</u>	<u>\$ 7,855</u>

The provision for income taxes included the following (in millions):

	Years ended December 31,		
	2025	2024	2023
Current provision:			
Federal	\$ 959	\$ 965	\$ 1,524
State	35	33	43
Foreign	971	759	786
Total current provision	<u>1,965</u>	<u>1,757</u>	<u>2,353</u>
Deferred benefit:			
Federal	(465)	(860)	(1,124)
State	7	(18)	(25)
Foreign	(242)	(360)	(66)
Total deferred benefit	<u>(700)</u>	<u>(1,238)</u>	<u>(1,215)</u>
Total provision for income taxes	<u>\$ 1,265</u>	<u>\$ 519</u>	<u>\$ 1,138</u>

Deferred income taxes reflect the tax effect of temporary differences between the carrying amounts of assets and liabilities for financial reporting purposes and the amounts used for income tax purposes, tax credit carryforwards and the tax effects of NOL carryforwards. As of December 31, 2022, we elected to establish deferred taxes with respect to the U.S. tax on the earnings of our foreign subsidiaries for the reversal of temporary items in future years. Significant components of our deferred tax assets and liabilities were as follows (in millions):

	December 31,	
	2025	2024
Deferred income tax assets:		
NOL and credit carryforwards	\$ 1,368	\$ 1,352
Accrued expenses	890	693
Capitalized research and development expenses	1,655	1,762
Investments	—	1
Expenses capitalized for tax	218	200
Earnings of foreign subsidiaries	2,305	1,496
Stock-based compensation	138	130
Other	349	361
Total deferred income tax assets	6,923	5,995
Valuation allowance	(1,299)	(1,019)
Net deferred income tax assets	5,624	4,976
Deferred income tax liabilities:		
Acquired intangible assets	(2,065)	(2,573)
Debt	(260)	(264)
Fixed assets	(192)	(143)
Fair value of acquired inventory	(28)	(114)
Investments	(422)	—
Other	(271)	(244)
Total deferred income tax liabilities	(3,238)	(3,338)
Total deferred income taxes, net	\$ 2,386	\$ 1,638

The Company has determined that unremitted foreign earnings are not considered indefinitely reinvested to the extent foreign earnings can be distributed without a significant tax cost.

Valuation allowances are provided to reduce the amounts of our deferred tax assets to an amount that is more likely than not to be realized based on an assessment of positive and negative evidence, including estimates of future taxable income necessary to realize future deductible amounts.

The valuation allowance increased in 2025, primarily driven by the Company's expectation that certain foreign amortization deductions are expected to expire unutilized.

As of December 31, 2025, we had \$109 million of federal tax credit carryforwards available to reduce future federal income taxes and have provided a \$31 million valuation allowance on those federal tax credit carryforwards. The federal tax credit carryforwards expire between 2032 and 2041. We had \$1.3 billion of state tax credit carryforwards available to reduce future state income taxes and have provided a valuation allowance for \$1.2 billion of those state tax credit carryforwards. We had \$148 million of tax credit carryforwards related to our foreign jurisdictions available to offset future foreign income taxes for which we have provided a \$123 million valuation allowance.

As of December 31, 2025, we had \$34 million of federal NOL carryforwards available to reduce future federal income taxes and have provided no valuation allowance on those federal NOL carryforwards. Additionally, \$12 million of those federal NOL carryforwards have no expiration; the remainder begin to expire between 2027 and 2033. We had \$971 million of state NOL carryforwards available to reduce future state income taxes and have provided a valuation allowance for \$836 million of those state NOL carryforwards. We had \$1.4 billion of foreign NOL carryforwards available to reduce future foreign income

taxes and have provided a valuation allowance for \$239 million of those foreign NOL carryforwards. For the foreign NOLs with no valuation allowance provided, \$120 million have no expiration; and the remainder will expire between 2026 and 2034.

The reconciliations of the total gross amounts of UTBs were as follows (in millions):

	Years ended December 31,		
	2025	2024	2023
Beginning balance	\$ 4,184	\$ 4,012	\$ 3,770
Additions based on tax positions related to the current year	190	188	196
Additions based on tax positions related to prior years	14	9	56
Reductions for tax positions of prior years	(13)	(12)	—
Reductions for expiration of statute of limitations	—	(9)	(4)
Settlements	(4)	(4)	(6)
Ending balance	<u>\$ 4,371</u>	<u>\$ 4,184</u>	<u>\$ 4,012</u>

Substantially all of the UTBs as of December 31, 2025, if recognized, would affect our effective tax rate. As a result, we remeasured our UTBs accordingly.

Interest and penalties related to UTBs are included in our provision for income taxes. During the years ended December 31, 2025, 2024 and 2023, we recognized \$264 million, \$282 million and \$287 million, respectively, of interest and penalties through the income tax provision in the Consolidated Statements of Income. The decrease in interest expense for the year ended December 31, 2025, was primarily due to lower interest rates in 2025 compared to those in 2024. As of December 31, 2025 and 2024, accrued interest and penalties associated with UTBs were \$1.9 billion and \$1.6 billion, respectively.

The reconciliations between the federal statutory tax rate applied to income before income taxes and our effective tax rate for 2025 were as follows (in millions, except percentages):

	Year ended December 31, 2025	
	Tax Effect	Effective Tax Rate
Federal statutory tax rate	\$ 1,885	21.0 %
Foreign tax effects:		
Ireland:		
Tax rate differential	316	3.5 %
Valuation allowance	119	1.3 %
Other	22	0.2 %
Puerto Rico:		
Tax rate differential	469	5.2 %
Act 52 - withholding tax	456	5.1 %
Act 52 - income tax	(968)	(10.8)%
Singapore:		
Tax incentive grant	(122)	(1.4)%
Pillar Two minimum tax	147	1.6 %
Other	(29)	(0.3)%
Other foreign jurisdictions	199	2.2 %
Effect of cross-border tax laws:		
U.S. tax on foreign subsidiaries (net of credits and deferred taxes)	(545)	(6.0)%
Foreign-derived intangible income	(222)	(2.5)%
Other	(18)	(0.2)%
Tax credits:		
Credit on foreign taxes imposed on U.S. income	(456)	(5.0)%
Research and development tax credits	(272)	(3.0)%
Other adjustments	86	1.0 %
Changes in unrecognized tax benefits (primarily interest)	198	2.2 %
Effective tax rate	\$ 1,265	14.1 %

The reconciliations between the federal statutory tax rate applied to income before income taxes and our effective tax rate for 2024 and 2023 were as follows:

	Years ended December 31,	
	2024	2023
Federal statutory tax rate	21.0 %	21.0 %
Foreign earnings	(5.8)%	(5.1)%
Foreign-derived intangible income	(3.0)%	(1.3)%
Credits, Puerto Rico excise tax	— %	0.3 %
Interest on uncertain tax positions	4.2 %	2.6 %
Credits, primarily federal R&D	(5.4)%	(3.5)%
Other, net	0.3 %	0.5 %
Effective tax rate	11.3 %	14.5 %

The effective tax rates for the years ended December 31, 2025, 2024 and 2023, differ from the federal statutory rate primarily due to impacts of the jurisdictional mix of income and expenses. Substantially all of the benefit to our effective tax rate from foreign earnings results from locations in which the Company has significant manufacturing operations, including Singapore, Ireland and Puerto Rico, a territory of the United States that is treated as a foreign jurisdiction for U.S. tax purposes. Our operations in Puerto Rico are subject to tax incentive grants through 2050 and the Company's operations in Singapore are

subject to a tax incentive grant through 2036. Effective January 1, 2024, selected individual countries, including the United Kingdom and EU member countries, have enacted the global minimum tax agreement. Additional countries, including Singapore, enacted the minimum tax agreement effective January 1, 2025. Singapore's enactment of the agreement applies irrespective of the Company's incentive grant. Due to the currently enacted scope of the agreement, the Company and its subsidiaries are now subject to a 15% minimum tax rate on adjusted financial statement income. Our foreign earnings are also subject to U.S. tax at a reduced rate of 10.5%.

Income taxes paid (net of refunds received) included the following (in millions):

	Year ended December 31, 2025
U.S. payments (federal and state)	\$ 3,390
Puerto Rico payments	610
Other foreign payments	300
Total income taxes paid (net of refunds received)	<u>\$ 4,300</u>

Income taxes paid (net of refunds received) during the years ended December 31, 2024 and 2023, were \$2.9 billion and \$3.4 billion, respectively.

On July 4, 2025, OB3 was enacted in the United States. OB3 has various provisions, including the permanent extension of certain expiring provisions of the 2017 Tax Act, and modifications to the international tax framework. The legislation has multiple effective dates, with certain provisions effective in 2026 and beyond. The impact of these changes was recorded in 2025.

One or more of our legal entities file income tax returns in the U.S. federal jurisdiction, various U.S. state jurisdictions and certain foreign jurisdictions. Our income tax returns are routinely examined by tax authorities in those jurisdictions. Significant disputes can arise and have arisen with tax authorities involving issues regarding the timing and amount of deductions, the use of tax credits and allocations of income and expenses among various tax jurisdictions because of differing interpretations of tax laws, regulations and relevant facts. Tax authorities, including the IRS, are becoming more aggressive and are particularly focused on such matters.

In 2017, we received an RAR and a modified RAR from the IRS for the years 2010–2012, proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico. We disagreed with the proposed adjustments and calculations, and in 2021, filed a petition in the U.S. Tax Court to contest two duplicate Statutory Notices of Deficiency (Notices) for the years 2010–2012. The Notices seek to increase our U.S. taxable income for the years 2010–2012 by an amount that would result in additional federal tax of approximately \$3.6 billion plus interest. Any additional tax that could be imposed for the years 2010–2012 would be reduced by up to approximately \$900 million of repatriation tax previously accrued and paid on our foreign earnings.

In 2020, we received an RAR and a modified RAR from the IRS for the years 2013–2015, also proposing significant adjustments that primarily relate to the allocation of profits between certain of our entities in the United States and the U.S. territory of Puerto Rico similar to those proposed for the years 2010–2012. We disagreed with the proposed adjustments and calculations, and in 2022, filed a petition in the U.S. Tax Court to contest a Notice for the years 2013–2015. The Notice seeks to increase our U.S. taxable income for the years 2013–2015 by an amount that would result in additional federal tax of approximately \$5.1 billion, plus interest, and asserts penalties of approximately \$2.0 billion. Any additional tax that could be imposed for the years 2013–2015 would be reduced by up to approximately \$2.2 billion of repatriation tax previously accrued and paid on our foreign earnings.

We firmly believe that the IRS positions set forth in the 2010–2012 and 2013–2015 Notices are without merit. We continue to contest the 2010–2012 and 2013–2015 Notices through the judicial process. The two cases were consolidated in the U.S. Tax Court in 2022. The trial began on November 4, 2024 and concluded on January 17, 2025. The parties filed opening post-trial briefs on June 13, 2025, and the Court held oral argument on July 16, 2025. The parties filed post-trial reply briefs on October 10, 2025. The Company expects a decision from the U.S. Tax Court no earlier than the second half of 2026.

We are currently under examination by the IRS for the years 2016–2018 with respect to issues similar to those for the 2010 through 2015 period. We expect that the IRS will begin its audit of 2019–2022 in the first half of 2026, and we believe that it may seek to continue to audit similar issues related to the allocation of income between the United States and our foreign jurisdictions. In addition, we are under examination by a number of state and foreign tax jurisdictions.

Final resolution of these complex matters is not likely within the next 12 months. We continue to believe our accrual for income tax liabilities is appropriate based on past experience, interpretations of tax law, application of the tax law to our facts

and judgments about potential actions by tax authorities; however, due to the complexity of the provision for income taxes and uncertain resolution of these matters, the ultimate outcome of any tax matters may result in payments substantially greater than amounts accrued and could have a material adverse impact on our consolidated financial statements.

We are no longer subject to U.S. federal income tax examinations for years ended on or before December 31, 2009.

8. Earnings per share

The computation of basic EPS is based on the weighted-average number of our common shares outstanding. The computation of diluted EPS is based on the weighted-average number of our common shares outstanding and dilutive potential common shares, which primarily include shares that may be issued under our stock option, restricted stock and performance unit award programs (collectively, dilutive securities), as determined by using the treasury stock method.

The computations for basic and diluted earnings per share were as follows (in millions, except per-share data):

	Years ended December 31,		
	2025	2024	2023
Income (Numerator):			
Net income for basic and diluted earnings per share	\$ 7,711	\$ 4,090	\$ 6,717
Shares (Denominator):			
Weighted-average shares for basic earnings per share	538	537	535
Effect of dilutive securities	4	4	3
Weighted-average shares for diluted earnings per share	542	541	538
Basic earnings per share			
	\$ 14.33	\$ 7.62	\$ 12.56
Diluted earnings per share			
	\$ 14.23	\$ 7.56	\$ 12.49

For each of the three years ended December 31, 2025, the number of antidilutive employee stock-based awards excluded from the computation of diluted EPS was not significant.

9. Collaborations

A collaborative arrangement is a contractual arrangement that involves a joint operating activity. Such arrangements involve two or more parties that are both (i) active participants in the activity and (ii) exposed to significant risks and rewards dependent on the commercial success of the activity.

From time to time, we enter into collaborative arrangements for the R&D, manufacture and/or commercialization of products and/or product candidates. These collaborations generally provide for nonrefundable upfront license fees, development and commercial-performance milestone payments, cost sharing, royalties and/or profit sharing. Our collaboration arrangements are performed with no guarantee of either technological or commercial success, and each arrangement is unique in nature. See Note 1, Summary of significant accounting policies, for additional discussion of revenues recognized under these types of arrangements. Operating expenses for costs incurred pursuant to these arrangements are reported in their respective expense line items in the Consolidated Statements of Income, net of any payments due to or reimbursements due from our collaboration partners, with such reimbursements being recognized at the time the party becomes obligated to pay. Our significant arrangements are discussed below.

AstraZeneca plc

We are in a collaboration with AstraZeneca for the development and commercialization of TEZSPIRE. Under our collaboration, both companies share global costs, profits and losses equally after payment by AstraZeneca of a mid-single-digit royalty to Amgen. AstraZeneca leads global development. In North America, Amgen, as the principal, recognizes product sales of TEZSPIRE in the United States, and AstraZeneca, as the principal, recognizes product sales of TEZSPIRE in Canada. AstraZeneca leads commercialization for TEZSPIRE outside North America. Amgen manufactures and supplies TEZSPIRE worldwide.

During the years ended December 31, 2025, 2024 and 2023, global profit and loss share expenses were \$587 million, \$412 million and \$310 million, respectively, and were recorded primarily in Cost of sales in the Consolidated Statements of

Income. Net costs due to AstraZeneca for global development and commercialization were not material during the years ended December 31, 2025, 2024 and 2023.

UCB

We are in a collaboration with UCB for the development and commercialization of EVENITY. Under our collaboration, UCB has rights to lead commercialization for EVENITY in most countries in Europe. Amgen, as the principal, leads commercialization for EVENITY and recognizes product sales in all other territories, including the United States. Global development costs and commercialization profits and losses related to the collaboration are shared equally. Amgen manufactures and supplies EVENITY worldwide.

During the years ended December 31, 2025, 2024 and 2023, global profit and loss share expenses were \$721 million, \$547 million and \$396 million, respectively, and were recorded primarily in Cost of sales in the Consolidated Statements of Income. Net costs recovered from and due to UCB during the years ended December 31, 2025, 2024 and 2023, were not material.

BeOne Medicines Ltd.

In January 2020, we acquired an equity stake in BeOne for approximately \$2.8 billion in cash as part of a collaboration agreement to expand our oncology presence in China. For additional information regarding our equity investment in BeOne, see Note 10, Investments. Under the collaboration, BeOne began selling XGEVA in 2020, BLINCYTO in 2021 and KYPROLIS in 2022 in China, and Amgen shares profits and losses equally during the product-specific commercialization periods. Following entry into an amendment by Amgen and BeOne on November 11, 2025, BeOne will retain the right to commercialize XGEVA, BLINCYTO and KYPROLIS in China for so long as such products are sold in China. Amgen manufactures and supplies the collaboration products to BeOne.

In addition, we jointly develop a portion of our oncology portfolio with BeOne, which shares in global R&D costs by providing cash and development services of up to \$1.25 billion. Upon regulatory approval, BeOne will assume commercialization rights in China for a specified period, and Amgen and BeOne will share profits and losses equally until certain of these product rights revert to Amgen. Upon return of the product rights, Amgen will pay royalties to BeOne on sales in China for a specified period. For product sales outside China, Amgen also pays royalties to BeOne.

During the years ended December 31, 2025, 2024 and 2023, net costs recovered from BeOne for oncology product candidates were \$181 million, \$122 million and \$109 million, respectively, and were recorded as an offset to R&D expense in the Consolidated Statements of Income. During the years ended December 31, 2025, 2024 and 2023, product sales from Amgen to BeOne under the collaboration were \$266 million, \$259 million and \$125 million, respectively, and were recorded in Product sales in the Consolidated Statements of Income. Profit and loss share expenses related to the product-specific commercialization period were not material during the years ended December 31, 2025, 2024 and 2023. Royalties on product sales of oncology portfolio products outside of China were not material for the years ended December 31, 2025, 2024 and 2023.

Kyowa Kirin Co., Ltd.

Since 2021, we have been in a collaboration agreement with Kyowa Kirin to jointly develop and commercialize rocatinlimab, an anti-OX40 fully human monoclonal antibody, worldwide, except in Japan. Rocatinlimab is for the treatment of atopic dermatitis, with potential for treatment of other autoimmune diseases.

Amgen and Kyowa Kirin share equally the global development costs, except in Japan, and the U.S. commercialization costs. Outside the United States and Japan, any commercialization costs incurred by Kyowa Kirin will be reimbursed by Amgen. We may also be required to make milestone payments of up to \$850 million contingent upon the achievement of certain regulatory events and commercial thresholds. We will also pay Kyowa Kirin significant double-digit royalties on global sales, except in Japan.

In January 2026, we and Kyowa Kirin agreed to terminate the rocatinlimab collaboration agreement and to transition control of the global development and commercialization program to Kyowa Kirin, subject to receipt of clearance under the Hart-Scott-Rodino Antitrust Improvements Act of 1976. In February 2026, we received such regulatory clearance. In the first quarter of 2026, Kyowa Kirin will assume full responsibility for rocatinlimab worldwide, except that Amgen will continue to manufacture rocatinlimab and perform other transition activities for an agreed upon period of time.

During the years ended December 31, 2025, 2024 and 2023, net costs recovered from Kyowa Kirin were \$117 million, \$166 million and \$93 million, respectively, and were recorded as an offset to R&D expense in the Consolidated Statements of Income.

Other

In addition to the collaborations discussed above, we have various other collaborations that are not individually significant to our business at this time. Pursuant to the terms of those agreements, we may be required to pay additional amounts, or we may receive additional amounts upon the achievement of various development and commercial milestones that in the aggregate could be significant. We may also incur or have reimbursed to us significant R&D costs if a related product candidate were to advance to late-stage clinical trials. In addition, if any products related to these collaborations are approved for sale, we may be required to pay significant royalties, or we may receive significant royalties on future sales. The payments of these amounts, however, are contingent upon the occurrence of various future events that have high degrees of uncertainty of occurrence.

10. Investments

Available-for-sale investments

The amortized cost, gross unrealized gains, gross unrealized losses and fair values of interest-bearing securities, which are classified as available for sale, by type of security were as follows (in millions):

Types of securities as of December 31, 2025	Amortized cost	Gross unrealized gains	Gross unrealized losses	Fair values
U.S. Treasury bills	\$ 998	\$ —	\$ —	\$ 998
Money market mutual funds	7,395	—	—	7,395
Other short-term interest-bearing securities	132	—	—	132
Total available-for-sale investments	<u>\$ 8,525</u>	<u>\$ —</u>	<u>\$ —</u>	<u>\$ 8,525</u>

Types of securities as of December 31, 2024	Amortized cost	Gross unrealized gains	Gross unrealized losses	Fair values
U.S. Treasury bills	\$ 997	\$ —	\$ —	\$ 997
Money market mutual funds	10,354	—	—	10,354
Other short-term interest-bearing securities	135	—	—	135
Total available-for-sale investments	<u>\$ 11,486</u>	<u>\$ —</u>	<u>\$ —</u>	<u>\$ 11,486</u>

The fair values of available-for-sale investments by location in the Consolidated Balance Sheets were as follows (in millions):

Consolidated Balance Sheets locations	December 31,	
	2025	2024
Cash and cash equivalents	\$ 8,525	\$ 11,486
Total available-for-sale investments	<u>\$ 8,525</u>	<u>\$ 11,486</u>

Cash and cash equivalents in the above table excludes bank account cash of \$604 million and \$487 million as of December 31, 2025 and 2024, respectively.

All interest-bearing securities as of December 31, 2025 and 2024, mature in one year or less. For the years ended December 31, 2025, 2024 and 2023, interest income on these investments was \$408 million, \$510 million and \$1.2 billion, respectively.

For the years ended December 31, 2025, 2024 and 2023, realized gains and losses on interest-bearing securities were not material and were recorded in Other income, net, in the Consolidated Statements of Income. The cost of securities sold is based on the specific-identification method.

The primary objective of our investment portfolio is to maintain safety of principal, prudent levels of liquidity and acceptable levels of risk. Our investment policy limits interest-bearing security investments to certain types of debt and money market instruments issued by institutions with investment-grade credit ratings, and it places restrictions on maturities and concentration by asset class and issuer.

Equity securities

BeOne Medicines Ltd.

As of December 31, 2025 and 2024, our ownership interest in BeOne was approximately 17.1% and 17.8%, respectively, and the fair values of our investment were \$5.8 billion and \$3.5 billion, respectively, which were included in Other noncurrent assets in the Consolidated Balance Sheets. We account for our ownership interest as an equity security with a readily determinable fair value, which is carried at fair value with changes in fair value recorded in Other income, net, in the Consolidated Statements of Income. See Note 18, Fair value measurement. During the years ended December 31, 2025, 2024 and 2023, we recognized unrealized gains of \$2.3 billion, \$82 million, and \$1.2 billion respectively, in Other income, net, in the Consolidated Statements of Income. For information on a collaboration agreement we entered into with BeOne in connection with this investment, see Note 9, Collaborations.

Subject to certain exceptions or otherwise agreed to by BeOne, while Amgen holds at least 5.0% of BeOne's outstanding common stock, (A) we may only sell our BeOne equity investment via: (i) a registered public offering, (ii) a sale under Rule 144 of the Securities Act of 1933 (the "Securities Act") or (iii) a private sale exempt from registration requirements under the Securities Act, and (B) we may not sell more than 5.0% of BeOne's outstanding common stock in any rolling 12-month period.

Other equity securities

Excluding our equity investments in BeOne (discussed above) and Neumora (discussed below), we held investments in other equity securities with readily determinable fair values (publicly traded securities) of \$326 million and \$314 million as of December 31, 2025 and 2024, respectively, which were included in Other noncurrent assets in the Consolidated Balance Sheets. For the years ended December 31, 2025, 2024 and 2023, net unrealized gains and losses on publicly traded securities resulted in a net gain of \$109 million, a net loss of \$21 million and a net gain of \$98 million, respectively. Realized gains and losses on publicly traded securities for the years ended December 31, 2025, 2024 and 2023, were not material.

We held investments of \$362 million and \$319 million in equity securities without readily determinable fair values as of December 31, 2025 and 2024, respectively, which were included in Other noncurrent assets in the Consolidated Balance Sheets. For the years ended December 31, 2025, 2024 and 2023, gains due to upward adjustments and gains realized upon dispositions of these securities were not material. For the years ended December 31, 2025, 2024 and 2023 downward adjustments were not material. Adjustments were based on observable price transactions.

Equity Method Investments

Neumora Therapeutics, Inc.

As of December 31, 2025 and 2024, our ownership interest in Neumora was approximately 21.2% and 21.9%, respectively, and the fair values of our investment were \$63 million and \$375 million, respectively, which were included in Other noncurrent assets in the Consolidated Balance Sheets. Although our equity investment qualifies us for the equity method of accounting, we have elected the fair value option to account for our investment. See Note 18, Fair value measurement. Under the fair value option, changes in the fair value of the investment are recognized through earnings in Other income, net, in the Consolidated Statements of Income each reporting period. We believe the fair value option best reflects the economics of the underlying transaction. During the years ended December 31, 2025, 2024 and 2023, we recognized unrealized losses of \$312 million and \$228 million and an unrealized gain of \$238 million, respectively, for the change in fair values in Other income, net, in the Consolidated Statements of Income.

We are contractually restricted from selling more than 5.0% of Neumora's outstanding common stock in any rolling 12-month period for as long as we hold at least 10.0% of their outstanding common stock, subject to certain exceptions or otherwise agreed to by Neumora.

Limited partnerships

We held limited partnership investments of \$253 million and \$262 million as of December 31, 2025 and 2024, respectively, which were included in Other noncurrent assets in the Consolidated Balance Sheets. These investments, which are primarily investment funds of early-stage biotechnology companies, are accounted for by using the equity method of accounting and are measured by using our proportionate share of the net asset values of the underlying investments held by the limited partnerships as a practical expedient. These investments are typically redeemable only through distributions upon liquidation of the underlying assets. As of December 31, 2025, we had \$134 million of unfunded additional commitments to be made for these investments during the next several years. For the years ended December 31, 2025, 2024 and 2023, net gains and losses recognized from our limited partnership investments were not material.

11. Inventories

Inventories consisted of the following (in millions):

	December 31,	
	2025	2024
Raw materials	\$ 915	\$ 818
Work in process	3,425	4,120
Finished goods	1,885	2,060
Total inventories	<u>\$ 6,225</u>	<u>\$ 6,998</u>

12. Property, plant and equipment

Property, plant and equipment consisted of the following (dollar amounts in millions):

	Useful life (in years)	December 31,	
		2025	2024
Land	—	\$ 348	\$ 346
Buildings and improvements	10-40	4,932	4,803
Manufacturing equipment	8-12	3,589	3,291
Laboratory equipment	8-12	1,438	1,345
Fixed equipment	12	2,668	2,592
Capitalized software	3-5	1,554	1,442
Other	5-10	1,114	1,059
Construction in progress	—	3,390	2,053
Property, plant and equipment, gross		<u>19,033</u>	<u>16,931</u>
Less accumulated depreciation and amortization		<u>(11,120)</u>	<u>(10,388)</u>
Property, plant and equipment, net		<u>\$ 7,913</u>	<u>\$ 6,543</u>

During the years ended December 31, 2025, 2024 and 2023, we recognized depreciation and amortization expense associated with our property, plant and equipment of \$763 million, \$694 million and \$685 million, respectively.

Geographic information

Certain geographic information with respect to property, plant and equipment, net, was as follows (in millions):

	December 31,	
	2025	2024
U.S.	\$ 5,188	\$ 4,156
Puerto Rico	1,273	1,174
ROW	1,452	1,213
Total property, plant and equipment, net	<u>\$ 7,913</u>	<u>\$ 6,543</u>

13. Goodwill and other intangible assets

Goodwill

The changes in the carrying amounts of goodwill were as follows (in millions):

	December 31,	
	2025	2024
Beginning balance	\$ 18,637	\$ 18,629
Changes to goodwill resulting from acquisitions, net ⁽¹⁾	—	25
Foreign currency translation adjustments	43	(17)
Ending balance	\$ 18,680	\$ 18,637

⁽¹⁾ For 2024, changes to Goodwill consisted of measurement-period adjustments related to our Horizon acquisition.

Other intangible assets

Other intangible assets consisted of the following (in millions):

	December 31,					
	2025			2024		
	Gross carrying amounts	Accumulated amortization	Other intangible assets, net	Gross carrying amounts	Accumulated amortization	Other intangible assets, net
Finite-lived intangible assets:						
Developed-product-technology rights	\$ 47,805	\$ (26,754)	\$ 21,051	\$ 48,611	\$ (22,594)	\$ 26,017
Licensing rights	3,917	(3,522)	395	3,875	(3,392)	483
R&D technology rights	1,425	(1,305)	120	1,374	(1,235)	139
Marketing-related rights	1,203	(1,203)	—	1,202	(1,202)	—
Total finite-lived intangible assets	54,350	(32,784)	21,566	55,062	(28,423)	26,639
Indefinite-lived intangible assets:						
In-process research and development	710	—	710	1,060	—	1,060
Total other intangible assets	\$ 55,060	\$ (32,784)	\$ 22,276	\$ 56,122	\$ (28,423)	\$ 27,699

Developed-product-technology rights consists of rights related to marketed products acquired in business combinations. Licensing rights primarily consists of contractual rights to receive future milestone, royalty and profit-sharing payments; capitalized payments to third parties for milestones related to regulatory approvals to commercialize products; and upfront payments associated with royalty obligations for marketed products. R&D technology rights pertains to technologies used in R&D that have alternative future uses. Marketing-related rights primarily consists of rights related to the sale and distribution of marketed products.

The Company monitors intangible assets for impairment on a quarterly basis. In January 2025, as part of the IRA, the Company's product Otezla was selected by CMS for Medicare price setting that will be applicable beginning on January 1, 2027. The earlier than anticipated selection resulted in a decrease in the estimated future cash flows for the product in the United States. This selection represented a triggering event that required the Company to evaluate the underlying developed-product-technology rights for impairment. In the first quarter of 2025, the Company utilized a discounted cash flow analysis based on Level 3 inputs, including estimated product sales, operating expenses and a discount rate, that resulted in an intangible asset fair value of \$4.0 billion, which was lower than the carrying value of \$4.8 billion, and a partial impairment of \$800 million. In the third quarter of 2025, new facts and circumstances, primarily from the CMS price setting process, indicated a further triggering event that required the Company to evaluate the underlying developed-product-technology rights for impairment. A subsequent discounted cash flow analysis, prepared using the same Level 3 input framework and updated assumptions, resulted in a revised intangible asset fair value of \$3.0 billion, which was lower than the carrying value of \$3.4 billion, and an additional impairment of \$400 million. The aggregate impairment charges of \$1.2 billion during the year ended December 31, 2025 were recognized in Other operating expenses in the Consolidated Statements of Income. See Note 18, Fair value measurement.

The developed-product-technology rights intangible assets related to TAVNEOS have a carrying value of \$2.5 billion as of December 31, 2025 with \$2.4 billion related to the U.S. market. On January 16, 2026, the FDA requested that ChemoCentryx voluntarily withdraw TAVNEOS from the U.S. market. Amgen is confident that TAVNEOS demonstrates effectiveness and a favorable benefit–risk profile. On January 28, 2026, following FDA regulatory process, Amgen informed the FDA that it did not intend to withdraw TAVNEOS from the market. Amgen is evaluating next steps with the FDA to determine a path forward. Future changes to estimated TAVNEOS cash flows could unfavorably impact the Company’s ability to recover the carrying value of the related intangible asset.

IPR&D consists of R&D projects acquired in a business combination that are not complete at the time of acquisition due to remaining technological risks and/or lack of receipt of required regulatory approvals. All IPR&D projects have major risks and uncertainties associated with the timely and successful completion of the development and commercialization of product candidates, including our ability to confirm safety and efficacy based on data from clinical trials, our ability to obtain necessary regulatory approvals and our ability to successfully complete these tasks within budgeted costs. We are not permitted to market a human therapeutic without obtaining regulatory approvals, and such approvals require the completion of clinical trials that demonstrate that a product candidate is safe and effective. In addition, the availability and extent of coverage and reimbursement from third-party payers, including government healthcare programs and private insurance plans as well as competitive product launches, affect the revenues a product can generate. Consequently, the eventual realized values, if any, of acquired IPR&D projects may vary from their estimated fair values. We review IPR&D projects for impairment annually, whenever events or changes in circumstances indicate that the carrying amounts may not be recoverable and upon the establishment of technological feasibility or regulatory approval. During the year ended December 31, 2023, the development of AMG 340 acquired in connection with our Teneobio acquisition was terminated, resulting in an impairment charge of \$783 million, which was recognized in Other operating expenses in the Consolidated Statements of Income.

During the second quarter of 2025, the FDA approved UPLIZNA for the IgG4-RD indication, and commercialization commenced in the United States. As a result, the Company reclassified the related intangible asset with a gross carrying value of \$350 million from IPR&D to developed-product-technology rights and began amortizing it on a straight-line basis over its estimated useful life of approximately 11 years from the date placed in service.

During the years ended December 31, 2025, 2024 and 2023, we recognized amortization of our finite-lived intangible assets of \$4.3 billion, \$4.8 billion and \$3.2 billion, respectively. Amortization of intangible assets is primarily included in Cost of sales in the Consolidated Statements of Income. As of December 31, 2025, the total estimated future amortization of our finite-lived intangible assets for the years ending December 31, 2026, 2027, 2028, 2029 and 2030, was \$3.6 billion, \$3.5 billion, \$2.8 billion, \$2.3 billion and \$2.2 billion, respectively.

14. Leases

We lease certain facilities and equipment related primarily to R&D, administrative and commercial activities. Leases with terms of 12 months or less are expensed as incurred and are not recorded in the Consolidated Balance Sheets.

Most leases include one or more options to renew, with renewal terms that may extend the lease term up to ten years. The exercise of lease renewal options is at our sole discretion. In addition, some of our lease agreements include rental payments adjusted periodically for inflation. Our lease agreements neither contain residual value guarantees nor impose significant restrictions or covenants. We sublease certain real estate to third parties. Our sublease portfolio consists of operating leases from former R&D and administrative spaces.

The following table summarizes information related to our leases, all of which are classified as operating, included in our Consolidated Balance Sheets (in millions):

Consolidated Balance Sheets locations	December 31,	
	2025	2024
Assets:		
Other noncurrent assets	\$ 602	\$ 557
Liabilities:		
Accrued liabilities	\$ 135	\$ 107
Other noncurrent liabilities	696	673
Total lease liabilities	\$ 831	\$ 780

The components of net lease costs were as follows (in millions):

Lease costs	Years ended December 31,		
	2025	2024	2023
Operating ⁽¹⁾	\$ 265	\$ 219	\$ 208
Sublease income	(9)	(17)	(28)
Total net lease costs	\$ 256	\$ 202	\$ 180

⁽¹⁾ Includes short-term leases and variable lease costs, which were not material for the years ended December 31, 2025, 2024 and 2023.

Maturities of lease liabilities as of December 31, 2025, were as follows (in millions):

Maturity dates	Amounts
2026	\$ 148
2027	153
2028	128
2029	99
2030	77
Thereafter	362
Total lease payments ⁽¹⁾	967
Less imputed interest	(136)
Present value of lease liabilities	\$ 831

⁽¹⁾ Includes future rental commitments for properties that have been subleased in the amount of \$99 million. We expect to receive total future rental income of \$54 million related to noncancellable subleases.

The weighted-average remaining lease terms and weighted-average discount rates were as follows:

	December 31,	
	2025	2024
Weighted-average remaining lease term (in years)	8.7	9.3
Weighted-average discount rate	4.0 %	3.7 %

Cash and noncash information related to our leases was as follows (in millions):

	Years ended December 31,		
	2025	2024	2023
Cash paid for amounts included in the measurement of lease liabilities:			
Operating cash flows for operating leases	\$ 157	\$ 156	\$ 182
ROU assets obtained in exchange for lease obligations:			
Operating leases	\$ 217	\$ 126	\$ 245

As of December 31, 2025, there were no future lease payments for leases that have not yet commenced.

15. Other current assets and accrued liabilities

Other current assets consisted of the following (in millions):

	December 31,	
	2025	2024
Prepaid expenses	\$ 2,945	\$ 2,139
Other	1,188	1,138
Total other current assets	\$ 4,133	\$ 3,277

Accrued liabilities consisted of the following (in millions):

	December 31,	
	2025	2024
Sales deductions	\$ 10,606	\$ 8,405
Employee compensation and benefits	1,437	1,329
Dividends payable	1,358	1,278
Income taxes payable	379	2,583
Other	4,743	4,046
Total accrued liabilities	\$ 18,523	\$ 17,641

16. Financing arrangements

Our borrowings consisted of the following (in millions):

	December 31,	
	2025	2024
1.90% notes due 2025 (1.90% 2025 Notes)	\$ —	\$ 500
5.25% notes due 2025 (5.25% 2025 Notes)	—	2,000
3.125% notes due 2025 (3.125% 2025 Notes)	—	1,000
2.00% €750 million notes due 2026 (2.00% 2026 euro Notes)	881	777
5.507% notes due 2026 (5.507% 2026 Notes)	—	1,500
2.60% notes due 2026 (2.60% 2026 Notes)	1,250	1,250
Term loan due October 2026	1,800	1,800
5.50% £475 million notes due 2026 (5.50% 2026 pound sterling Notes)	640	595
2.20% notes due 2027 (2.20% 2027 Notes)	1,724	1,724
3.20% notes due 2027 (3.20% 2027 Notes)	1,000	1,000
5.15% notes due 2028 (5.15% 2028 Notes)	3,750	3,750
1.65% notes due in 2028 (1.65% 2028 Notes)	1,234	1,234
3.00% notes due 2029 (3.00% 2029 Notes)	750	750
4.05% notes due 2029 (4.05% 2029 Notes)	1,250	1,250
4.00% £700 million notes due 2029 (4.00% 2029 pound sterling Notes)	944	876
2.45% notes due 2030 (2.45% 2030 Notes)	1,250	1,250
5.25% notes due 2030 (5.25% 2030 Notes)	2,750	2,750
2.30% notes due 2031 (2.30% 2031 Notes)	1,250	1,250
2.00% notes due 2032 (2.00% 2032 Notes)	987	1,001
3.35% notes due 2032 (3.35% 2032 Notes)	1,000	1,000
4.20% notes due 2033 (4.20% 2033 Notes)	750	750
5.25% notes due 2033 (5.25% 2033 Notes)	4,250	4,250
6.375% notes due 2037 (6.375% 2037 Notes)	478	478
6.90% notes due 2038 (6.90% 2038 Notes)	254	254
6.40% notes due 2039 (6.40% 2039 Notes)	333	333
3.15% notes due 2040 (3.15% 2040 Notes)	1,478	1,668
5.75% notes due 2040 (5.75% 2040 Notes)	373	373
2.80% notes due 2041 (2.80% 2041 Notes)	568	776
4.95% notes due 2041 (4.95% 2041 Notes)	600	600
5.15% notes due 2041 (5.15% 2041 Notes)	729	729
5.65% notes due 2042 (5.65% 2042 Notes)	415	415
5.60% notes due 2043 (5.60% 2043 Notes)	2,750	2,750
5.375% notes due 2043 (5.375% 2043 Notes)	185	185
4.40% notes due 2045 (4.40% 2045 Notes)	2,250	2,250
4.563% notes due 2048 (4.563% 2048 Notes)	1,415	1,415
3.375% notes due 2050 (3.375% 2050 Notes)	1,462	1,764
4.663% notes due 2051 (4.663% 2051 Notes)	3,541	3,541
3.00% notes due 2052 (3.00% 2052 Notes)	703	890
4.20% notes due 2052 (4.20% 2052 Notes)	882	895
4.875% notes due 2053 (4.875% 2053 Notes)	1,000	1,000
5.65% notes due 2053 (5.65% 2053 Notes)	4,250	4,250
2.77% notes due 2053 (2.77% 2053 Notes)	940	940
4.40% notes due 2062 (4.40% 2062 Notes)	1,128	1,165
5.75% notes due 2063 (5.75% 2063 Notes)	2,750	2,750

	December 31,	
	2025	2024
Other notes due 2097	100	100
Total principal amount of debt	56,044	61,778
Unamortized bond discounts, premiums and issuance costs, net	(1,306)	(1,360)
Fair value adjustments	(161)	(343)
Other	27	24
Total carrying value of debt	54,604	60,099
Less current portion	(4,599)	(3,550)
Total long-term debt	\$ 50,005	\$ 56,549

There are no material differences between the effective interest rates and coupon rates of our notes, except for the 4.563% 2048 Notes, the 4.663% 2051 Notes and the 2.77% 2053 Notes, which have effective interest rates of 6.3%, 5.6% and 5.2%, respectively.

Under the terms of all of our outstanding notes, except our Other notes due 2097, in the event of a change-in-control triggering event we may be required to purchase all or a portion of these debt securities at prices equal to 101% of the principal amounts of the notes plus accrued and unpaid interest. In addition, all of our outstanding notes—except our Other notes due 2097—may be redeemed at any time at our option—in whole or in part—at the principal amounts of the notes being redeemed plus accrued and unpaid interest and make-whole amounts, which are defined by the terms of the notes. Certain of the redeemable notes do not require the payment of make-whole amounts if redeemed during a specified period of time immediately prior to the maturity of the notes. Such time periods range from one month to six months prior to maturity.

Debt issuances and acquisition-related financing

We did not issue debt securities during the years ended December 31, 2025 and 2024.

In March 2023, in connection with the acquisition of Horizon (see Note 4, Acquisition), we issued the following series of notes (in millions):

	Principal Amount
5.25% 2025 Notes	\$ 2,000
5.507% 2026 Notes	1,500
5.15% 2028 Notes	3,750
5.25% 2030 Notes	2,750
5.25% 2033 Notes	4,250
5.60% 2043 Notes	2,750
5.65% 2053 Notes	4,250
5.75% 2063 Notes	2,750
Total	\$ 24,000

Also in connection with the acquisition of Horizon, we entered into a \$4.0 billion term loan credit agreement in December 2022. In October 2023, in connection with the completion of the acquisition of Horizon, we borrowed \$4.0 billion under the term loan credit agreement, of which \$2.2 billion was repaid during 2024. As of December 31, 2025, we had \$1.8 billion of borrowings outstanding under the term loan credit agreement, which has an interest rate of three-month SOFR plus 1.225% and is due in October 2026.

Debt extinguishment

In 2025, we repurchased an aggregate principal amount of our debt of \$1.0 billion, including portions of the 2.00% 2032 Notes, 3.15% 2040 Notes, 2.80% 2041 Notes, 3.375% 2050 Notes, 3.00% 2052 Notes, 4.20% 2052 Notes and 4.40% 2062 Notes, for an aggregate cost of \$683 million, which resulted in a \$264 million gain on extinguishment of debt recorded in Other income, net, in the Consolidated Statements of Income.

In 2024, we repurchased an aggregate principal amount of our debt of \$875 million, including portions of the 3.15% 2040 Notes, 2.80% 2041 Notes, 3.375% 2050 Notes, 3.00% 2052 Notes, 4.20% 2052 Notes and 4.40% 2062 Notes, for an aggregate

cost of \$659 million, which resulted in a \$215 million gain on extinguishment of debt recorded in Other income, net, in the Consolidated Statements of Income.

In 2023, we repurchased an aggregate principal amount of our debt of \$881 million, including portions of the 2.00% 2032 Notes, 3.15% 2040 Notes, 2.80% 2041 Notes, 3.375% 2050 Notes, 3.00% 2052 Notes, 4.20% 2052 Notes and 4.40% 2062 Notes, for an aggregate cost of \$647 million, which resulted in a \$225 million gain on extinguishment of debt recorded in Other income, net, in the Consolidated Statements of Income.

Debt repayments

In 2025, debt repayments totaled \$5.0 billion, including repayments in full of the \$500 million aggregate principal amount of the 1.90% 2025 Notes, \$2.0 billion aggregate principal amount of the 5.25% 2025 Notes, \$1.0 billion aggregate principal amount of the 3.125% 2025 Notes and \$1.5 billion aggregate principal amount of the 5.507% 2026 Notes.

In 2024, debt repayments totaled \$3.6 billion, including the full \$2.0 billion aggregate principal amount on the term loan due April 2025, \$200 million of the aggregate principal amount on the term loan due October 2026 and the full \$1.4 billion aggregate principal amount of the 3.625% 2024 Notes.

In 2023, debt repayments totaled \$1.5 billion, including the full \$750 million aggregate principal amount of the 2.25% 2023 Notes and the full CHF700 million aggregate principal amount (\$704 million upon settlement of the related cross-currency swap) of the 0.41% 2023 Swiss franc Bonds.

Interest rate swaps

To achieve a desired mix of fixed-rate and floating-rate debt, we enter into interest rate swap contracts that effectively convert fixed-rate interest coupons for certain of our debt instruments to floating SOFR-based coupons over the terms of the respective debt instruments. These interest rate swap contracts qualify and are designated as fair value hedges. For information regarding the terms of these contracts, see Note 19, Derivative instruments.

Cross-currency swaps

To hedge our exposure to foreign currency exchange rate risk associated with certain of our long-term notes denominated in foreign currencies, we entered into cross-currency swap contracts. The terms of these contracts outstanding as of December 31, 2025, effectively convert the interest payments and principal repayments on our 2.00% 2026 euro Notes, 5.50% 2026 pound sterling Notes and 4.00% 2029 pound sterling Notes from euros and pounds sterling to U.S. dollars. These cross-currency swap contracts have been designated as cash flow hedges. For information regarding the terms of these contracts, see Note 19, Derivative instruments. Cross-currency swap contracts associated with other foreign denominated debt previously outstanding were settled in connection with the repayment of such debt, as discussed above.

Shelf registration statement and other facilities

As of December 31, 2025, we have a commercial paper program that allows us to issue up to \$4.0 billion of unsecured commercial paper to fund working capital needs. As of December 31, 2025 and 2024, we had no amounts outstanding under our commercial paper program.

In the first quarter of 2023, we amended and restated our syndicated, unsecured, revolving credit agreement, under which we may borrow up to \$4.0 billion for general corporate purposes, including as a liquidity backstop for our commercial paper program. The commitments under the revolving credit agreement may be increased by up to \$1.25 billion with the agreement of the banks. Each bank that is a party to the agreement has an initial commitment term of five years. This term may be extended for up to two additional one-year periods with the agreement of the banks. Annual commitment fees for this agreement are 0.09% of the unused portion of the facility based on our current credit rating. Generally, we would be charged interest for any amounts borrowed under this facility, based on our current credit rating, at (i) SOFR plus 1.01% or (ii) the highest of (A) the administrative agent bank base commercial lending rate, (B) the overnight federal funds rate plus 0.50% or (C) one-month SOFR plus 1.1%. As of December 31, 2025 and 2024, no amounts were outstanding under this facility.

In February 2023, we filed a shelf registration statement with the SEC that allows us to issue unspecified amounts of debt securities; common stock; preferred stock; warrants to purchase debt securities, common stock, preferred stock or depositary shares; rights to purchase common stock or preferred stock; securities purchase contracts; securities purchase units; and depositary shares. Under this shelf registration statement, all of the securities available for issuance may be offered from time to time, with terms to be determined at the time of issuance. This shelf registration statement expired in February 2026, and our Board has approved a new shelf registration statement to replace it.

Certain of our financing arrangements contain nonfinancial covenants. In addition, our revolving credit agreement and term loan agreement include a financial covenant, which requires us to maintain a specified minimum interest coverage ratio of (i) the sum of consolidated net income, interest expense, provision for income taxes, depreciation expense, amortization expense, unusual or nonrecurring charges and other noncash items (Consolidated EBITDA) to (ii) Consolidated Interest Expense, each as defined and described in the respective agreements. We were in compliance with all applicable covenants under these arrangements as of December 31, 2025.

Contractual maturities of debt obligations

The aggregate contractual maturities of our debt obligations as of December 31, 2025, were as follows (in millions):

Maturity dates	Amounts
2026	\$ 4,571
2027	2,724
2028	4,984
2029	2,943
2030	4,000
Thereafter	36,822
Total	\$ 56,044

Interest costs

Interest costs are expensed as incurred except to the extent such interest is related to construction in progress, in which case interest is capitalized. Interest costs capitalized for the years ended December 31, 2025, 2024 and 2023, were not material. Interest paid, net of amounts capitalized and including the ongoing impact of interest rate and cross-currency swap contracts, during the years ended December 31, 2025, 2024 and 2023 was \$2.9 billion, \$3.3 billion and \$2.4 billion, respectively.

17. Stockholders' equity

Stock repurchase program

During the years ended December 31, 2025 and 2023, we did not repurchase shares under our stock repurchase program. During the year ended December 31, 2024, we repurchased 0.7 million shares of our common stock for a total cost of \$200 million under our stock repurchase program. As of December 31, 2025, \$6.8 billion of authorization remained available under the stock repurchase program.

Dividends

Our Board of Directors declared quarterly dividends per share of \$2.38, \$2.25 and \$2.13, which were paid in each of the four quarters of 2025, 2024 and 2023, respectively.

Historically, we have declared dividends in December of each year, which were paid in the first quarter of the following fiscal year and in March, July and October, which were paid in the second, third and fourth quarters, respectively, of the same fiscal year. Additionally, in December 2025, the Board of Directors declared a quarterly cash dividend of \$2.52 per share of common stock, which will be paid in March 2026, to all stockholders of record as of the close of business on February, 13 2026.

Accumulated other comprehensive loss

The components of AOCI were as follows (in millions):

	Foreign currency translation adjustments	Cash flow hedges	Other	AOCI
Balance as of December 31, 2022	\$ (348)	\$ 128	\$ (11)	\$ (231)
Foreign currency translation adjustments	50	—	—	50
Unrealized gains	—	28	—	28
Reclassification adjustments into earnings	—	(222)	—	(222)
Other	—	—	42	42
Income taxes	—	44	—	44
Balance as of December 31, 2023	(298)	(22)	31	(289)
Foreign currency translation adjustments	(76)	—	—	(76)
Unrealized gains	—	506	—	506
Reclassification adjustments into earnings	—	(117)	—	(117)
Other	—	—	(10)	(10)
Income taxes	—	(80)	—	(80)
Balance as of December 31, 2024	(374)	287	21	(66)
Foreign currency translation adjustments	172	—	—	172
Unrealized losses	—	(250)	—	(250)
Reclassification adjustments into earnings	—	(231)	—	(231)
Other	—	—	13	13
Income taxes	—	104	—	104
Balance as of December 31, 2025	<u>\$ (202)</u>	<u>\$ (90)</u>	<u>\$ 34</u>	<u>\$ (258)</u>

With respect to the table above, income tax expenses or benefits for unrealized gains and losses and the related reclassification adjustments to earnings for cash flow hedges were a \$53 million benefit and a \$51 million benefit in 2025, a \$105 million expense and a \$25 million benefit in 2024, and a \$6 million expense and a \$50 million benefit in 2023, respectively.

Reclassifications out of AOCI and into earnings were as follows (in millions):

Components of AOCI	Years ended December 31,			Consolidated Statements of Income locations
	2025	2024	2023	
Cash flow hedges:				
Foreign currency forward contract gains	\$ 10	\$ 192	\$ 180	Product sales
Cross-currency swap contract gains (losses)	221	(75)	42	Other income, net
	231	117	222	Income before income taxes
	(51)	(25)	(50)	Provision for income taxes
	<u>\$ 180</u>	<u>\$ 92</u>	<u>\$ 172</u>	Net income

Other

In addition to common stock, our authorized capital includes 5 million shares of preferred stock, \$0.0001 par value. As of December 31, 2025 and 2024, no shares of preferred stock were issued or outstanding.

18. Fair value measurement

To estimate the fair values of our financial assets and liabilities, we use valuation approaches within a hierarchy that maximizes the use of observable inputs and minimizes the use of unobservable inputs by requiring that observable inputs be used when available. Observable inputs are inputs that market participants would use in pricing an asset or liability based on market data obtained from sources independent of the Company. Unobservable inputs are inputs that reflect the Company's assumptions about the inputs that market participants would use in pricing an asset or liability and are developed based on the best information available in the circumstances. The fair value hierarchy is divided into three levels based on the source of inputs as follows:

- Level 1 — Valuations based on unadjusted quoted prices in active markets for identical assets or liabilities that the Company has the ability to access
- Level 2 — Valuations for which all significant inputs are observable either directly or indirectly—other than Level 1 inputs
- Level 3 — Valuations based on inputs that are unobservable and significant to the overall fair value measurement

The availability of observable inputs can vary among the various types of financial assets and liabilities. To the extent that the valuation is based on models or inputs that are less observable or unobservable in the market, the determination of fair value requires more judgment. In certain cases, the inputs used for measuring fair value may fall into different levels of the fair value hierarchy. In such cases, for financial statement disclosure purposes, the level in the fair value hierarchy within which the fair value measurement is categorized is based on the lowest level of input used that is significant to the overall fair value measurement.

The fair values of each major class of the Company's financial assets and liabilities measured at fair value on a recurring basis were as follows (in millions):

Fair value measurement as of December 31, 2025, using:	Quoted prices in active markets for identical assets (Level 1)	Significant other observable inputs (Level 2)	Significant unobservable inputs (Level 3)	Total
Assets:				
Available-for-sale securities:				
U.S. Treasury bills	\$ —	\$ 998	\$ —	\$ 998
Money market mutual funds	7,395	—	—	7,395
Other short-term interest-bearing securities	—	132	—	132
Equity securities	6,144	—	—	6,144
Derivatives:				
Foreign currency forward contracts	—	196	—	196
Cross-currency swap contracts	—	48	—	48
Total assets	<u>\$ 13,539</u>	<u>\$ 1,374</u>	<u>\$ —</u>	<u>\$ 14,913</u>
Liabilities:				
Derivatives:				
Foreign currency forward contracts	\$ —	\$ 214	\$ —	\$ 214
Cross-currency swap contracts	—	320	—	320
Interest rate swap contracts	—	293	—	293
Contingent consideration obligations	—	—	161	161
Total liabilities	<u>\$ —</u>	<u>\$ 827</u>	<u>\$ 161</u>	<u>\$ 988</u>

Fair value measurement as of December 31, 2024, using:	Quoted prices in active markets for identical assets (Level 1)	Significant other observable inputs (Level 2)	Significant unobservable inputs (Level 3)	Total
Assets:				
Available-for-sale securities:				
U.S. Treasury bills	\$ —	\$ 997	\$ —	\$ 997
Money market mutual funds	10,354	—	—	10,354
Other short-term interest-bearing securities	—	135	—	135
Equity securities	4,188	—	—	4,188
Derivatives:				
Foreign currency forward contracts	—	420	—	420
Total assets	<u>\$ 14,542</u>	<u>\$ 1,552</u>	<u>\$ —</u>	<u>\$ 16,094</u>
Liabilities:				
Derivatives:				
Foreign currency forward contracts	\$ —	\$ 8	\$ —	\$ 8
Cross-currency swap contracts	—	483	—	483
Interest rate swap contracts	—	531	—	531
Contingent consideration obligations	—	—	106	106
Total liabilities	<u>\$ —</u>	<u>\$ 1,022</u>	<u>\$ 106</u>	<u>\$ 1,128</u>

Interest-bearing and equity securities

The fair values of our U.S. Treasury bills are determined by utilizing third-party pricing services, which obtain pricing data from active market makers and brokers. The fair values of our money market mutual funds and equity investments in publicly traded securities, including our equity investments in BeOne and Neumora, as of December 31, 2025 and 2024, are based on quoted market prices in active markets, with no valuation adjustment.

Derivatives

All of our foreign currency forward contracts, cross-currency swap contracts and interest rate swap contracts are with counterparties that have minimum credit ratings of A– or equivalent by S&P, Moody's or Fitch. We estimate the fair values of these contracts by taking into consideration valuations obtained from a third-party valuation service that uses an income-based industry-standard valuation model for which all significant inputs are observable either directly or indirectly. These inputs, as applicable, include foreign currency exchange rates, SOFR, swap rates, obligor credit default swap rates and cross-currency basis swap spreads. Certain inputs, when applicable, are at commonly quoted intervals. See Note 19, Derivative instruments.

Contingent consideration obligations

As a result of business development activity, we have incurred contingent consideration obligations as discussed below. The contingent consideration obligations are recorded at their fair values by using probability-adjusted discounted cash flows, and we revalue these obligations each reporting period until the related contingencies have been resolved. The fair value measurements of these obligations are based on significant unobservable inputs related to licensing rights and product candidates acquired through business development activity, and they are reviewed quarterly by management in our R&D and commercial sales organizations. The inputs include, as applicable, estimated probabilities and the timing of achieving specified development, regulatory and commercial milestones as well as estimated annual sales. Significant changes that increase or decrease the probabilities of achieving the related development, regulatory and commercial events or that shorten or lengthen the time required to achieve such events or that increase or decrease estimated annual sales would result in corresponding increases or decreases in the fair values of the obligations, as applicable. Changes in the fair values of contingent consideration obligations are recognized in Other operating expenses in the Consolidated Statements of Income.

Changes in the carrying amounts of contingent consideration obligations were as follows (in millions):

	Years ended December 31,	
	2025	2024
Beginning balance	\$ 106	\$ 96
Additions	68	—
Payments	(7)	(8)
Net changes in valuations	(6)	18
Ending balance	\$ 161	\$ 106

As of December 31, 2025 and 2024, our contingent consideration obligations were primarily the result of our acquisition of Teneobio in October 2021, which obligates us to make payments to the former shareholders upon achievement of separate development and regulatory milestones with regard to various R&D programs, and other business development activity in 2025.

Summary of the fair values of other financial instruments

Cash equivalents

The fair values of cash equivalents are approximated at their carrying values due to the short-term nature of such financial instruments.

Borrowings

We estimate the fair values of our fixed-rate debt by using Level 2 inputs. As of December 31, 2025 and 2024, the aggregate fair values of our fixed-rate debt were \$51.0 billion and \$54.9 billion, respectively, and the carrying values of our fixed-rate debt were \$52.8 billion and \$58.3 billion, respectively. The estimate of the fair value of our term loan is approximated at its carrying value as of December 31, 2025 and 2024, as this debt instrument bears interest at a floating rate.

During the years ended December 31, 2025 and 2024, there were no transfers of assets or liabilities between fair value measurement levels. Except with respect to the partial impairments of the Otezla intangible asset in 2025 and the IPR&D intangible impairment of AMG 340 in 2023 as disclosed in Note 13, Goodwill and other intangible assets, there were no material remeasurements of the fair values of assets and liabilities that are not measured at fair value on a recurring basis.

19. Derivative instruments

The Company is exposed to foreign currency exchange rate and interest rate risks related to its business operations. To reduce our risks related to such exposures, we use or have used certain derivative instruments, including foreign currency forward, foreign currency option, cross-currency swap, forward interest rate and interest rate swap contracts. We have designated certain of our derivatives as cash flow and fair value hedges; we also have derivatives not designated as hedges. We do not use derivatives for speculative trading purposes.

Cash flow hedges

We are exposed to possible changes in the values of certain anticipated foreign currency cash flows resulting from changes in foreign currency exchange rates primarily associated with our euro-denominated international product sales. The foreign currency exchange rate fluctuation exposure associated with cash inflows from our international product sales is partially offset by corresponding cash outflows from our international operating expenses. To further reduce this exposure, we enter into foreign currency forward contracts to hedge a portion of our projected international product sales up to a maximum of three years into the future; and at any given point in time, a higher percentage of nearer-term projected product sales is being hedged than in successive periods.

As of December 31, 2025 and 2024, we had outstanding foreign currency forward contracts with aggregate notional amounts of \$7.8 billion and \$7.2 billion, respectively. We have designated these foreign currency forward contracts, which are primarily euro based, as cash flow hedges. Accordingly, we record unrealized gains and losses on these contracts in AOCI in the Consolidated Balance Sheets, and we reclassify them to Product sales in the Consolidated Statements of Income in the same periods during which the hedged transactions affect earnings.

To hedge our exposure to foreign currency exchange rate risk associated with certain of our long-term debt denominated in foreign currencies, we enter into cross-currency swap contracts. Under the terms of such contracts, we paid euros and pounds sterling and received U.S. dollars for the notional amounts at the inception of the contracts; and based on these notional amounts, we exchange interest payments at fixed rates over the terms of the contracts by paying U.S. dollars and receiving euros and pounds sterling. In addition, we will pay U.S. dollars to and receive euros and pounds sterling from the counterparties at the maturities of the contracts for these same notional amounts. The terms of these contracts correspond to the related hedged debt, thereby effectively converting the interest payments and principal repayment on the debt from euros and pounds sterling to U.S. dollars. We have designated these cross-currency swap contracts as cash flow hedges. Accordingly, the unrealized gains and losses on these contracts are recorded in AOCI in the Consolidated Balance Sheets and reclassified to Other income, net, in the Consolidated Statements of Income in the same periods during which the hedged debt affects earnings.

The notional amounts and interest rates of our cross-currency swaps as of December 31, 2025, were as follows (notional amounts in millions):

Hedged notes	Foreign currency		U.S. dollars	
	Notional amounts	Interest rates	Notional amounts	Interest rates
2.00% 2026 euro Notes	€ 750	2.0 %	\$ 833	3.9 %
5.50% 2026 pound sterling Notes	£ 475	5.5 %	\$ 747	6.0 %
4.00% 2029 pound sterling Notes	£ 700	4.0 %	\$ 1,111	4.7 %

In connection with the anticipated issuance of long-term fixed-rate debt, we occasionally enter into forward interest rate contracts in order to hedge the variability in cash flows due to changes in the applicable U.S. Treasury rate between the time we enter into these contracts and the time the related debt is issued. Gains and losses on forward interest rate contracts, which are designated as cash flow hedges, are recognized in AOCI in the Consolidated Balance Sheets and are amortized into Interest expense, net, in the Consolidated Statements of Income over the terms of the associated debt issuances. In 2025, we entered into forward interest rate contracts with an aggregate notional amount of \$500 million. Amounts recognized in connection with forward interest rate contracts during the years ended December 31, 2025, 2024 and 2023, and amounts expected to be recognized during the next 12 months on forward interest rate contracts were not material.

Unrealized gains and losses recognized in AOCI for our derivative instruments designated as cash flow hedges were as follows (in millions):

Derivatives in cash flow hedging relationships	Years ended December 31,		
	2025	2024	2023
Foreign currency forward contracts	\$ (464)	\$ 585	\$ (14)
Cross-currency swap contracts	214	(79)	73
Forward interest rate contracts	—	—	(31)
Total unrealized (losses) gains	\$ (250)	\$ 506	\$ 28

Fair value hedges

To achieve a desired mix of fixed-rate and floating-rate debt, we enter into interest rate swap contracts that qualify for and are designated as fair value hedges. These interest rate swap contracts effectively convert fixed-rate coupons to floating-rate SOFR-based coupons over the terms of the related hedge contracts. As of both December 31, 2025 and 2024, we had interest rate swap contracts with an aggregate notional amount of \$6.7 billion that hedge certain portions of our long-term debt. In 2025, interest rate swap contracts with an aggregate notional amount of \$1.0 billion matured in connection with the repayment of the 3.125% 2025 Notes. Also in 2025, we entered into \$1.0 billion of new interest rate swap contracts to hedge a portion of our 5.25% 2033 Notes.

As of December 31, 2025 and 2024, the interest rates on the portion of notes for which we have entered into interest rate swap contracts and the related notional amounts of these contracts were as follows (dollar amounts in millions):

Notes	December 31,			
	2025		2024	
	Notional amounts	Interest rates	Notional amounts	Interest rates
3.125% 2025 Notes	\$ —	N/A	\$ 1,000	SOFR + 2.1%
2.60% 2026 Notes	1,250	SOFR + 2.1%	1,250	SOFR + 2.1%
2.45% 2030 Notes	1,000	SOFR + 1.3%	1,000	SOFR + 1.3%
2.30% 2031 Notes	500	SOFR + 1.1%	500	SOFR + 1.1%
5.25% 2033 Notes	2,400	SOFR + 1.8%	1,400	SOFR + 1.8%
4.663% 2051 Notes	1,500	SOFR + 4.3%	1,500	SOFR + 4.3%
Total notional amounts	<u>\$ 6,650</u>		<u>\$ 6,650</u>	

N/A = not applicable

For interest rate swap contracts that qualify for and are designated as fair value hedges, we recognize in Interest expense, net, in the Consolidated Statements of Income the unrealized gain or loss on the derivative resulting from the change in fair value during the period, as well as the offsetting unrealized loss or gain of the hedged item resulting from the change in fair value during the period attributable to the hedged risk. If a hedging relationship involving an interest rate swap contract is terminated, the gain or loss realized on contract termination is recorded as an adjustment to the carrying value of the debt and amortized into Interest expense, net, over the remaining term of the previously hedged debt.

The hedged liabilities and related cumulative-basis adjustments for fair value hedges of those liabilities were recorded in the Consolidated Balance Sheets as follows (in millions):

Consolidated Balance Sheets locations	Carrying amounts of hedged liabilities ⁽¹⁾		Cumulative amounts of fair value hedging adjustments related to the carrying amounts of the hedged liabilities ⁽²⁾	
	December 31,		December 31,	
	2025	2024	2025	2024
Current portion of long-term debt	\$ 1,273	\$ 1,045	\$ 23	\$ 45
Long-term debt	\$ 5,112	\$ 5,152	\$ (184)	\$ (388)

⁽¹⁾ Current portion of long-term debt includes \$47 million and \$56 million of carrying value with discontinued hedging relationships as of December 31, 2025 and 2024, respectively. Long-term debt includes \$185 million and \$232 million of carrying value with discontinued hedging relationships as of December 31, 2025 and 2024, respectively.

⁽²⁾ Current portion of long-term debt includes \$47 million and \$56 million of hedging adjustments on discontinued hedging relationships as of December 31, 2025 and 2024, respectively. Long-term debt includes \$85 million and \$132 million of hedging adjustments on discontinued hedging relationships as of December 31, 2025 and 2024, respectively.

Impact of hedging transactions

The following tables summarize the amounts recorded in income and expense line items and the effects thereon from fair value and cash flow hedging, including discontinued hedging relationships (in millions):

	Year ended December 31, 2025		
	Product sales	Other income, net	Interest expense, net
Total amounts recorded in income and (expense) line items presented in the Consolidated Statements of Income	\$ 35,148	\$ 2,651	\$ (2,755)
The effects of cash flow and fair value hedging:			
Gains on cash flow hedging relationships reclassified out of AOCI:			
Foreign currency forward contracts	\$ 10	\$ —	\$ —
Cross-currency swap contracts	\$ —	\$ 221	\$ —
(Losses) gains on fair value hedging relationships—interest rate swap agreements:			
Hedged items ⁽¹⁾	\$ —	\$ —	\$ (182)
Derivatives designated as hedging instruments	\$ —	\$ —	\$ 238
	Year ended December 31, 2024		
	Product sales	Other income, net	Interest expense, net
Total amounts recorded in income and (expense) line items presented in the Consolidated Statements of Income	\$ 32,026	\$ 506	\$ (3,155)
The effects of cash flow and fair value hedging:			
Gains (losses) on cash flow hedging relationships reclassified out of AOCI:			
Foreign currency forward contracts	\$ 192	\$ —	\$ —
Cross-currency swap contracts	\$ —	\$ (75)	\$ —
Gains on fair value hedging relationships—interest rate swap agreements:			
Hedged items ⁽¹⁾	\$ —	\$ —	\$ 29
Derivatives designated as hedging instruments	\$ —	\$ —	\$ 40
	Year ended December 31, 2023		
	Product sales	Other income, net	Interest expense, net
Total amounts recorded in income and (expense) line items presented in the Consolidated Statements of Income	\$ 26,910	\$ 2,833	\$ (2,875)
The effects of cash flow and fair value hedging:			
Gains on cash flow hedging relationships reclassified out of AOCI:			
Foreign currency forward contracts	\$ 180	\$ —	\$ —
Cross-currency swap contracts	\$ —	\$ 42	\$ —
(Losses) gains on fair value hedging relationships—interest rate swap agreements:			
Hedged items ⁽¹⁾	\$ —	\$ —	\$ (118)
Derivatives designated as hedging instruments	\$ —	\$ —	\$ 205

⁽¹⁾ Gains (losses) on hedged items do not exactly offset losses (gains) on the related designated hedging instruments due to amortization of the cumulative amounts of fair value hedging adjustments included in the carrying amount of the hedged debt for discontinued hedging relationships and the recognition of gains on terminated hedges when the corresponding hedged item was paid down in the period.

No portions of our cash flow hedge contracts were excluded from the assessment of hedge effectiveness. As of December 31, 2025, the amount of net losses on our foreign currency forward and cross-currency swap contracts expected to be reclassified out of AOCI and recognized into earnings during the next 12 months was \$120 million.

Derivatives not designated as hedges

To reduce our exposure to foreign currency fluctuations in certain assets and liabilities denominated in foreign currencies, we enter into foreign currency forward contracts that are not designated as hedging transactions. Most of these exposures are hedged on a month-to-month basis. As of December 31, 2025 and 2024, the total notional amounts of these foreign currency forward contracts were \$240 million and \$148 million, respectively. Gains and losses recognized in earnings for our derivative instruments not designated as hedging instruments were not material for the years ended December 31, 2025, 2024 and 2023.

Fair values of derivatives

The fair values of derivatives included in the Consolidated Balance Sheets were as follows (in millions):

December 31, 2025	Derivative assets		Derivative liabilities	
	Consolidated Balance Sheets locations	Fair values	Consolidated Balance Sheets locations	Fair values
Derivatives designated as hedging instruments:				
Foreign currency forward contracts	Other current assets/ Other noncurrent assets	\$ 195	Accrued liabilities/ Other noncurrent liabilities	\$ 213
Cross-currency swap contracts	Other current assets/ Other noncurrent assets	48	Accrued liabilities/ Other noncurrent liabilities	320
Interest rate swap contracts	Other current assets/ Other noncurrent assets	—	Accrued liabilities/ Other noncurrent liabilities	293
Total derivatives designated as hedging instruments		243		826
Derivatives not designated as hedging instruments:				
Foreign currency forward contracts	Other current assets	1	Accrued liabilities	1
Total derivatives not designated as hedging instruments		1		1
Total derivatives		\$ 244		\$ 827

December 31, 2024	Derivative assets		Derivative liabilities	
	Consolidated Balance Sheets locations	Fair values	Consolidated Balance Sheets locations	Fair values
Derivatives designated as hedging instruments:				
Foreign currency forward contracts	Other current assets/ Other noncurrent assets	\$ 420	Accrued liabilities/ Other noncurrent liabilities	\$ 8
Cross-currency swap contracts	Other current assets/ Other noncurrent assets	—	Accrued liabilities/ Other noncurrent liabilities	483
Interest rate swap contracts	Other current assets/ Other noncurrent assets	—	Accrued liabilities/ Other noncurrent liabilities	531
Total derivatives designated as hedging instruments		420		1,022
Derivatives not designated as hedging instruments:				
Foreign currency forward contracts	Other current assets	—	Accrued liabilities	—
Total derivatives not designated as hedging instruments		—		—
Total derivatives		\$ 420		\$ 1,022

For additional information, see Note 18, Fair value measurement.

Our derivative contracts that were in liability positions as of December 31, 2025, contain certain credit-risk-related contingent provisions that would be triggered if (i) we were to undergo a change in control and (ii) our or the surviving entity's creditworthiness deteriorates, which is generally defined as having either a credit rating that is below investment grade or a materially weaker creditworthiness after the change in control. If these events were to occur, the counterparties would have the right, but not the obligation, to close the contracts under early-termination provisions. In such circumstances, the counterparties could request immediate settlement of these contracts for amounts that approximate the then current fair values of the contracts. In addition, our derivative contracts are not subject to any type of master netting arrangement, and amounts due either to or from a counterparty under the contracts may be offset against other amounts due either to or from the same counterparty only if an event of default or termination, as defined, were to occur.

The cash flow effects of our derivative contracts in the Consolidated Statements of Cash Flows are primarily included in Net cash provided by operating activities, except for the settlement of notional amounts of cross-currency swaps, which are included in Net cash (used in) provided by financing activities.

20. Contingencies and commitments

Contingencies

In the ordinary course of business, we are involved in various legal proceedings, government investigations and other matters that are complex in nature and have outcomes that are difficult to predict. See Part I, Item 1A. Risk Factors—*Our business may be affected by litigation and government investigations*. We describe our legal proceedings and other matters that are significant or that we believe could become significant in this footnote.

We record accruals for loss contingencies to the extent that we conclude it is probable that a liability has been incurred and the amount of the related loss can be reasonably estimated. We evaluate, on a quarterly basis, developments in legal proceedings and other matters that could cause an increase or decrease in the amount of the liability that has been accrued previously.

Our legal proceedings involve various aspects of our business and a variety of claims, some of which present novel factual allegations and/or unique legal theories. The outcomes of these proceedings are inherently uncertain and depend on a variety of factors, including the development of the factual record, judicial or administrative rulings, and, in certain cases, the outcome of appellate review. Further, certain of the matters pending against us are at earlier stages of the legal process, which in complex proceedings of the sort we face often extend for several years, and have not progressed sufficiently through discovery and/or the development of important factual information and legal issues to enable us to estimate. Accordingly, except for amounts accrued, in each of the matters described in this filing in which we could incur a liability, our opponents seek an award of a not-yet-estimable amount of damages or an amount that is not material. While it is not possible to accurately predict or determine the eventual outcomes of these matters, an adverse determination in one or more of these matters currently pending could have a material adverse effect on our consolidated results of operations, financial position or cash flows.

Certain recent developments concerning our legal proceedings and other matters are discussed below.

Repatha Patent Litigation

Patent Disputes in the International Region

We are involved in and expect future involvement in additional disputes regarding our PCSK9 patents in other jurisdictions and regions. This includes matters filed against us and that we have filed in Germany and Japan.

Germany

In February 2016, the EPO granted European Patent No. 2,215,124 (the EP'124 Patent) to Amgen. This patent describes and claims monoclonal antibodies to PCSK9 and methods of treatment and Sanofi filed an opposition to the patent in the EPO seeking to invalidate it. In November 2016, Sanofi-Aventis Deutschland GmbH, Sanofi-Aventis Groupe S.A. and Sanofi Winthrop Industrie S.A. filed a joint opposition against Amgen's patent, and each of Eli Lilly and Company, Regeneron Pharmaceuticals, Inc. (Regeneron) and Strawman Ltd. also filed oppositions to Amgen's patent. In November 2018, the EPO confirmed the validity of Amgen's EP'124 Patent, which was appealed to the Technical Board of Appeal (TBA). On October 29, 2020, the TBA upheld the validity of certain claims, including claims that protect Repatha, but ruled that broader claims encompassing PRALUENT® were invalid. As a result of the TBA's decision, any national litigations regarding infringement of the EP'124 Patent by PRALUENT® were terminated.

In Germany, Sanofi-Aventis Deutschland GmbH and Regeneron filed actions in the Regional Court of Munich seeking damages arising from the provisional enforcement of an injunction against PRALUENT® that was lifted after the TBA's October 2020 ruling on the EP'124 Patent. On May 8, 2024, the Regional Court of Munich issued a preliminary decision. The case is still pending, but further proceedings were stayed on December 9, 2025.

Also in Germany, on July 21, 2022, Sanofi Biotechnology SAS filed an action against Amgen GmbH and Amgen (Europe) B.V. before the Regional Court of Dusseldorf alleging that the marketing and sale of Repatha infringes European Patent No. 2,756,004 (the EP'004 Patent), which Sanofi Biotechnology SAS licensed from Regeneron. Sanofi Biotechnology SAS is seeking infringement damages and injunctive relief. On May 13, 2024, the Regional Court of Dusseldorf stayed the hearing on Sanofi Biotechnology SAS' infringement action pending the outcome of Amgen's nullity action against the EP'004 Patent before the German Federal Patent Court. On December 29, 2025, the Regional Court of Dusseldorf lifted the stay and scheduled an oral hearing on May 12, 2026 to address whether Amgen infringes the EP'004 Patent in Germany.

On August 3, 2023, Amgen GmbH filed a nullity action before the German Federal Patent Court seeking invalidation of Regeneron's EP'004 Patent. On November 25, 2025, the German Federal Patent Court upheld the validity of Regeneron's EP'004 Patent.

Unified Patent Court of the European Union

Actions concerning Amgen's European Patent 3,666,797 (the EP'797 Patent)

On June 1, 2023, Amgen filed an action before the Munich Local Division of the Unified Patent Court (UPC) against Sanofi-Aventis Deutschland GmbH, Sanofi-Aventis Groupe S.A., Sanofi Winthrop Industrie S.A. (collectively, Sanofi-Aventis), and Regeneron alleging that the importation, marketing, sale and use of PRALUENT® infringes Amgen's EP'797 Patent seeking an injunction and damages for past infringement. Regeneron filed counterclaims for revocation, but on February 5, 2024, the court transferred the counterclaims to the Central Division of the UPC that is presiding over Sanofi's revocation action. The Munich Local Division of the UPC scheduled the hearing on Amgen's EP'797 Patent infringement action to begin on October 16, 2024.

On June 29, 2023, the Central Division of the UPC served Amgen with an action that was filed by Sanofi-Aventis that seeks revocation of the EP'797 Patent. The Central Division of the UPC scheduled a hearing on the revocation action and on July 16, 2024, the Central Division of the UPC rendered its decision, concluding that the patent claims are invalid and revoked the EP'797 Patent. Subsequently, on July 29, 2024, the Munich Local Division of the UPC stayed Amgen's action against Sanofi-Aventis alleging that the importation, marketing, sale and use of PRALUENT® infringes the EP'797 Patent. On September 13, 2024, Amgen filed a statement of appeal with the Court of Appeals of the UPC to set aside the Central Division of the UPC's decision to revoke the EP'797 Patent. On August 12, 2025, the Court of Appeals of the UPC heard oral argument on Amgen's appeal seeking to set aside the Central Division of the UPC's decision to revoke Amgen's EP'797 Patent and on November 25, 2025, the Court of Appeals of the UPC issued a decision upholding the validity of Amgen's EP'797 Patent. On January 26, 2026, Sanofi and Regeneron filed an application for rehearing and sought an order from the Court of Appeals of the UPC that the lodging of its application for rehearing suspends the UPC's decision upholding the validity of the EP'797 Patent. On January 29, 2026, the Court of Appeals of the UPC issued an order giving Amgen the opportunity to respond to the request for rehearing by February 26, 2026. On February 5, 2026, Amgen filed its reply to Sanofi's request for suspension.

On December 30, 2025, following on the Court of Appeals of the UPC's decision finding the EP'797 Patent valid, the Munich Local Division of the UPC lifted the stay on Amgen's case alleging that PRALUENT® infringes Amgen's EP'797 Patent. The Court scheduled an interim conference to take place on June 26, 2026, and an oral argument to take place on November 19, 2026.

Actions concerning Regeneron's European Patent 3,536,712 (the EP'712 Patent)

On January 10, 2024, Sanofi Biotechnologies SAS (Sanofi Biotechnologies) and Regeneron filed an action (the UPC Action Concerning the EP'712 Patent) against Amgen Inc., Amgen Europe B.V., Amgen N.V., Amgen GmbH, Amgen B.V., Amgen SAS, and Amgen S.R.L before the Dusseldorf Local Division of the UPC, alleging infringement of Regeneron's EP'712 Patent, which Sanofi Biotechnology SAS licensed from Regeneron. Sanofi and Regeneron are seeking an injunction against the sale, marketing, use, importation, or storage of Repatha for certain specified uses in Belgium, France, Germany, Italy and the Netherlands. Amgen filed counterclaims for invalidity and non-infringement. On February 25, 2025, a hearing on the validity and infringement of the EP'712 Patent was held before the Dusseldorf Local Division of the UPC.

On May 13, 2025, the Dusseldorf Local Division of the UPC issued a decision that the EP'712 Patent is valid but not infringed by Amgen. On July 11, 2025, Amgen filed its statement of appeal against the part of the Dusseldorf Local Division of the UPC's decision finding that the EP'712 Patent is valid. On July 14, 2025, Sanofi and Regeneron filed their statement of appeal against the decision of the Dusseldorf Local Division of the UPC finding the EP'712 Patent not infringed by Amgen. On September 15, 2025, Amgen and Sanofi and Regeneron filed their respective grounds of appeal. On December 15, 2025, the parties filed responses to the grounds of appeal.

Actions concerning Regeneron's European Patent 4,252,857 (the EP'857 Patent)

On September 25, 2024, Sanofi Biotechnologies and Regeneron filed a brief seeking to expand the ongoing action before the Dusseldorf Local Division of the UPC concerning the EP'712 Patent, alleging that Amgen's Repatha infringes a newly-issued patent, European Patent No. 4,252,857 (the EP'857 Patent) and seeking an injunction against the marketing, use, or importation of Repatha in 18 countries (Austria, Belgium, Bulgaria, Denmark, Estonia, Finland, France, Germany, Italy, Latvia, Lithuania, Luxembourg, Malta, the Netherlands, Portugal, Romania, Slovenia and Sweden) and damages for past infringement (the UPC Action Concerning the EP'857 Patent). On December 13, 2024, the Dusseldorf Local Division of the UPC denied Sanofi and Regeneron's request to extend the complaint. On February 4, 2025, the Dusseldorf Local Division of the UPC formally ordered separation of the EP'857 Patent from the ongoing EP'712 Patent litigation.

On June 9, 2025, Sanofi filed a motion seeking to stay the UPC Action Concerning the EP'857 Patent, which Amgen opposed. On August 6, 2025, the Dusseldorf Local Division of the UPC stayed the UPC Action Concerning the EP'857 Patent until the Court of Appeals of the UPC reaches a decision on the appeal from the UPC Action Concerning the EP'712 Patent.

European Patent Office

Proceedings concerning Amgen's EP'797 Patent

On November 16, 2023 and February 29, 2024, Sanofi-Aventis and Regeneron each filed a notice of opposition against Amgen's EP'797 Patent before the EPO's Opposition Division. A hearing was held beginning on March 31, 2025 and on April 3, 2025, the Opposition Division determined the claims of the EP'797 Patent are valid.

On April 16, 2025, Sanofi-Aventis and Regeneron filed notices of appeal and requested expedited appeal proceedings. The EPO TBA has scheduled a hearing on Sanofi's and Regeneron's appeals against the Opposition Division's decision upholding the validity of Amgen's EP'797 Patent to take place on April 13-15, 2026. On January 26, 2026, Amgen filed its reply to Sanofi-Aventis's and Regeneron's grounds of appeal.

Proceedings concerning Regeneron's EP'712 Patent

On February 29, 2024, Amgen filed a notice of opposition and grounds of opposition before the EPO against Regeneron's EP'712 Patent. On March 15, 2024, the EPO notified the parties that the opposition will be accelerated in view of the infringement action pending against Amgen on the EP'712 Patent in the Dusseldorf Local Division of the UPC. On March 12, 2025, following a hearing on Amgen's opposition to Regeneron's EP'712 Patent, the Opposition Division determined that the claims of the EP'712 Patent are valid, and issued its written decision on April 24, 2025.

On April 25, 2025, Amgen filed a notice of appeal and request for expedited appeal proceedings. On June 2, 2025, the EPO TBA accelerated Amgen's appeal from the EPO's decision that Regeneron's EP'712 Patent is valid and scheduled oral argument to take place on March 25, 2026.

Proceedings concerning Regeneron's EP'857 Patent

On June 23, 2025, Amgen filed a notice of opposition and grounds of opposition before the EPO against Regeneron's EP'857 Patent. On July 7, 2025, the EPO notified the parties that the opposition proceedings concerning the EP'857 Patent have been accelerated due to the pending parallel proceedings before the UPC, and Regeneron's response to Amgen's grounds of opposition must accordingly be filed by October 7, 2025. The EPO Opposition Division scheduled a hearing to take place on Amgen's opposition against Regeneron's EP'857 Patent on June 9-10, 2026.

Japan

On April 24, 2020, the Supreme Court of Japan declined to hear Sanofi K.K.'s appeals making final the Intellectual Property High Court's (IP High Court) decisions that PRALUENT® infringes Amgen's valid patent rights in Japan.

On June 24, 2020, Amgen filed written answers to the invalidity trials initiated by Regeneron on February 12, 2020 before the Japan Patent Office (JPO) seeking to invalidate Amgen's Japanese patents that were previously held infringed by PRALUENT® and valid over challenges filed by Sanofi K.K. On April 15, 2021, the JPO dismissed Regeneron's invalidity trials, and in August 2021 Regeneron appealed the decisions to the IP High Court. On January 26, 2023, the IP High Court found Amgen's patent claims invalid for lacking adequate support and Amgen appealed to the Supreme Court of Japan on March 13, 2023. On September 15, 2023, the Supreme Court of Japan declined to hear Amgen's appeal. The case was remanded to the JPO for further proceedings, however the case is currently stayed.

On March 31, 2021, Amgen initiated damages proceedings against Sanofi K.K. and Sanofi K.K. raised new validity challenges to Amgen patents. On September 27, 2023, the Tokyo District Court found Amgen's patent claims invalid and dismissed Amgen's lawsuit for damages. Amgen appealed the Tokyo District Court's decision to the IP High Court on December 28, 2023. The IP High Court rejected Amgen's appeal and remanded the case to the JPO. Amgen then sought amended patent claims before the JPO. The JPO rejected Amgen's amended claims, and Amgen appealed the JPO's decision to the IP High Court on September 16, 2024. On April 15, 2025, the IP High Court dismissed Amgen's appeal and, on May 27, 2025, Amgen filed petition for acceptance of an appeal before the Supreme Court of Japan.

Prolia/XGEVA Biologics Price Competition and Innovation Act (BPCIA) Litigation

Amgen Inc. et al. v. Shanghai Henlius Biotech Inc. et al.

On June 25, 2025, Amgen Inc. and Amgen Manufacturing Limited LLC filed a lawsuit in the U.S. District Court for the District of New Jersey (New Jersey District Court) against Shanghai Henlius Biotech Inc., Shanghai Henlius Biologics Co., Ltd,

Organon LLC and Organon & Co. (collectively, the Shanghai Henlius and Organon Defendants) based on the submission to the FDA of a BLA seeking approval to market and sell a biosimilar version of Amgen's Prolia and XGEVA products. The complaint asserts infringement of the following patents: U.S. Patent Nos. 7,364,736; 7,888,101; 7,928,205; 8,053,236; 8,217,153; 8,460,896; 8,680,248; 9,228,168; 9,359,435; 10,106,829; 10,227,627; 10,513,723; 10,583,397; 10,655,156; 10,894,972; 11,077,404; 11,098,079; 11,192,919; 11,254,963; 11,319,568; 11,434,514; 11,459,595; 11,492,372; 11,946,085; 11,952,605; and 12,084,686 (collectively, the Asserted Patents against the Shanghai Henlius and Organon Defendants). Amgen seeks a judgment from the New Jersey District Court that the Shanghai Henlius and Organon Defendants have infringed or will infringe one or more claims of each of the Asserted Patents against the Shanghai Henlius and Organon Defendants and based on that judgment, a permanent injunction prohibiting the commercial manufacture, use, offer to sell, or sale within the United States or importation into the United States of the accused proposed denosumab biosimilar by the Shanghai Henlius and Organon Defendants before expiration of each of the patents found to be infringed. Amgen also seeks monetary remedies for any past acts of infringement.

On June 25, 2025, this litigation became a member of In Re: Denosumab Patent Litigation multi-district litigation with other cases involving Prolia/XGEVA biosimilars pending in the district. On September 5, 2025, the Shanghai Henlius and Organon Defendants responded to the complaint, asserting counterclaims for invalidity and non-infringement and affirmative defenses. On October 10, 2025, Amgen responded to those counterclaims and asserted its affirmative defenses. A claim construction hearing will be scheduled after June 23, 2027, and the joint pretrial order is due April 28, 2028. A trial date has not yet been set.

Amgen Inc. et al. v. Hikma Pharmaceuticals USA Inc. et al.

On June 25, 2025, Amgen Inc. and Amgen Manufacturing Limited LLC filed a lawsuit in the New Jersey District Court against Hikma Pharmaceuticals USA Inc., Gedeon Richter Plc., and Gedeon Richter USA, Inc. (collectively, the Hikma and Gedeon Richter Defendants) based on the submission to the FDA of a BLA seeking approval to market and sell a biosimilar version of Amgen's Prolia and XGEVA products. The complaint asserts infringement of the following patents: U.S. Patent Nos. 7,364,736; 7,888,101; 7,928,205; 8,053,236; 8,058,418; 8,460,896; 8,680,248; 9,012,178; 9,228,168; 9,328,134; 9,359,435; 9,371,554; 10,106,829; 10,167,492; 10,227,627; 10,513,723; 10,583,397; 10,822,630; 10,894,972; 11,077,404; 11,098,079; 11,130,980; 11,192,919; 11,254,963; 11,299,760; 11,319,568; 11,434,514; 11,459,595; 11,492,372; 11,946,085; 11,952,605; and 12,084,686 (collectively, the Asserted Patents against the Hikma and Gedeon Richter Defendants). Amgen seeks a judgment from the New Jersey District Court that the Hikma and Gedeon Richter Defendants have infringed or will infringe one or more claims of each of the Asserted Patents against the Hikma and Gedeon Richter Defendants and based on that judgment, a permanent injunction prohibiting the commercial manufacture, use, offer to sell, or sale within the United States or importation into the United States of the accused proposed denosumab biosimilar before expiration of each of the patents found to be infringed. Amgen also seeks monetary remedies for any past acts of infringement. On June 25, 2025, this litigation became a member of In Re: Denosumab Patent Litigation multi-district litigation with other cases involving Prolia/XGEVA biosimilars pending in the district. A trial date has not yet been set.

On September 5, 2025, the Hikma and Gedeon Richter Defendants responded to the complaint, asserting counterclaims and affirmative defenses on September 5, 2025. On October 10, 2025, Amgen responded to those counterclaims and asserted its affirmative defenses. Pursuant to a consent order providing leave to amend, Amgen filed an Amended Complaint on November 14, 2025, adding Gedeon Richter (Schweiz) AG as a defendant to the litigation.

The parties entered into a settlement agreement that resolves the patent litigation related to the Hikma denosumab biosimilar products in the United States. Accordingly, the New Jersey District Court entered a Consent Judgment and Injunction on November 24, 2025, finding the claims of Amgen's U.S. patents asserted against the Hikma and Gedeon Richter Defendants valid, enforceable and infringed by Hikma's denosumab biosimilars in the United States and enjoining Defendants from importing, making, using, offering to sell or selling the Hikma denosumab biosimilar products in the United States before January 1, 2026, except as may be permitted by the agreement or by statute.

Amgen Inc. et al. v. Amneal Pharmaceuticals, Inc. et al.

On November 6, 2025, Amgen Inc. and Amgen Manufacturing Limited LLC filed a lawsuit in the New Jersey District Court against Amneal Pharmaceuticals, Inc., Amneal Pharmaceuticals LLC, GH Genhelix S.A., Universal Farma S.L., and Mabxience Research S.L., (collectively, the Amneal and Mabxience Defendants) based on the submission to the FDA of a BLA seeking approval to market and sell a biosimilar version of Amgen's Prolia and XGEVA products. The complaint asserts infringement of the following patents: U.S. Patent Nos. 7,364,736; 7,888,101; 7,928,205; 8,053,236; 8,058,418; 8,460,896; 8,680,248; 9,012,178; 9,228,168; 9,320,816; 9,328,134; 9,359,435; 10,106,829; 10,167,492; 10,227,627; 10,513,723; 10,583,397; 10,655,156; 10,822,630; 10,894,972; 11,077,404; 11,098,079; 11,130,980; 11,254,963; 11,299,760; 11,319,568; 11,434,514; 11,459,595; 11,946,085; 11,952,605; 12,084,686 (collectively, the Asserted Patents against the Amneal and Mabxience Defendants). Amgen seeks a judgment from the New Jersey District Court that the Amneal and Mabxience

Defendants have infringed or will infringe one or more claims of each of the Asserted Patents against the Amneal and Mabxience Defendants and based on that judgment, a permanent injunction prohibiting the commercial manufacture, use, offer to sell, or sale within the United States or importation into the United States of the accused denosumab biosimilar by the Amneal and Mabxience Defendants before expiration of each of the patents found to be infringed. Amgen also seeks monetary remedies for any past acts of infringement. On January 9, 2026 and January 23, 2026, respectively, the Amneal and Mabxience Defendants responded to the complaint asserting affirmative defenses.

This litigation is part of the In Re: Denosumab Patent Litigation multi-district litigation with other patent cases involving Prolia/XGEVA biosimilars pending in the district. A claim construction hearing will be scheduled after June 23, 2027, and the joint pretrial order is due April 28, 2028. A trial date has not yet been set.

Amgen Inc. et al. v. Dr. Reddy's Laboratories Ltd., et al.

On November 6, 2025, Amgen Inc. and Amgen Manufacturing Limited LLC filed a lawsuit in the New Jersey District Court against Dr. Reddy's Laboratories Ltd., Dr. Reddy's Laboratories SA, Dr. Reddy's Laboratories Inc., Alvotech Hf, and Alvotech Swiss AG (collectively, the Dr. Reddy's and Alvotech Defendants) based on the submission to the FDA of a BLA seeking approval to market and sell a biosimilar version of Amgen's Prolia and XGEVA products. The complaint asserts infringement of the following patents: U.S. Patent Nos. 7,364,736; 7,888,101; 7,928,205; 8,053,236; 8,058,418; 8,460,896; 8,680,248; 9,012,178; 9,228,168; 9,328,134; 9,359,435; 10,106,829; 10,167,492; 10,227,627; 10,513,723; 10,583,397; 10,822,630; 10,894,972; 11,077,404; 11,098,079; 11,130,980; 11,192,919; 11,254,963; 11,299,760; 11,319,568; 11,434,514; 11,459,595; 11,492,372; 11,946,085; 11,952,605; 12,084,686 (collectively, the Asserted Patents against the Dr. Reddy's and Alvotech Defendants). Amgen seeks a judgment from the New Jersey District Court that the Dr. Reddy's and Alvotech Defendants have infringed or will infringe one or more claims of each of the Asserted Patents against the Dr. Reddy's and Alvotech Defendants and based on that judgment, a permanent injunction prohibiting the commercial manufacture, use, offer to sell, or sale within the United States or importation into the United States of the accused proposed denosumab biosimilar by the Dr. Reddy's and Alvotech Defendants before expiration of each of the patents found to be infringed. Amgen also seeks monetary remedies for any past acts of infringement. On January 9, 2026, the Dr. Reddy's and Alvotech Defendants responded to the complaint asserting counterclaims for invalidity and non-infringement and affirmative defenses. On January 30, 2026, Amgen responded to those counterclaims and asserted its affirmative defenses.

This litigation is part of the In Re: Denosumab Patent Litigation multi-district litigation with other patent cases involving Prolia/XGEVA biosimilars pending in the district. A claim construction hearing will be scheduled after June 23, 2027, and the joint pretrial order is due April 28, 2028. A trial date has not yet been set.

Amgen Inc. et al. v. Alkem Laboratories Ltd., et al.

On November 14, 2025, Amgen Inc. and Amgen Manufacturing Limited LLC filed a lawsuit in the New Jersey District Court against Alkem Laboratories Ltd., Ascend Laboratories, LLC, and Enzene Biosciences (collectively, the Alkem Defendants) based on the submission to the FDA of a BLA seeking approval to market and sell a biosimilar version of Amgen's Prolia and XGEVA products. The complaint asserts infringement of the following patents: U.S. Patent Nos. 7,364,736; 7,662,930; 7,888,101; 7,928,205; 8,053,236; 8,058,418; 8,247,210; 8,460,896; 8,680,248; 9,012,178; 9,228,168; 9,328,134; 9,359,435; 10,106,829; 10,167,492; 10,227,627; 10,513,723; 10,583,397; 10,655,156; 10,822,630; 10,894,972; 10,907,186; 11,077,404; 11,098,079; 11,130,980; 11,192,919; 11,254,963; 11,299,760; 11,319,568; 11,434,514; 11,459,595; 11,492,372; 11,946,085; 11,952,605; 12,084,686 (collectively, the Asserted Patents against the Alkem Defendants). Amgen seeks a judgment from the New Jersey District Court that the Alkem Defendants have infringed or will infringe one or more claims of each of the Asserted Patents against the Alkem Defendants and based on that judgment, a permanent injunction prohibiting the commercial manufacture, use, offer to sell, or sale within the United States or importation into the United States of the accused proposed denosumab biosimilar by the Alkem Defendants before expiration of each of the patents found to be infringed. Amgen also seeks monetary remedies for any past acts of infringement. On January 23, 2026, the Alkem Defendants responded to the complaint asserting affirmative defenses and counterclaims. On February 12, 2026, the Alkem Defendants filed an amended answer asserting affirmative defenses and counterclaims.

This litigation is part of the In Re: Denosumab Patent Litigation multi-district litigation with other patent cases involving Prolia/XGEVA biosimilars pending in the district. A claim construction hearing will be scheduled after June 23, 2027, and the joint pretrial order is due April 28, 2028. A trial date has not yet been set.

PAVBLU® (afibercept-ayyh) Patent Litigation

On January 10, 2024, Regeneron filed a lawsuit in the U.S. District Court for the Central District of California (California Central District Court) against Amgen alleging infringement of 32 patents listed by Regeneron in the BPCIA exchange (the 2024 Complaint). The lawsuit stems from Amgen's submission of an application under the BPCIA for FDA licensure of

PAVBLU as biosimilar to Regeneron's EYLEA®. By its complaint, Regeneron seeks, among other remedies, an injunction prohibiting the commercial manufacture, use, offer for sale or sale in the United States or import into the United States of PAVBLU before the expiration of each of the patents found to be infringed. Amgen responded to Regeneron's complaint on February 2, 2024, denying infringement and asserting counterclaims seeking a declaratory judgment that the asserted patents are not infringed, invalid, and/or unenforceable.

On April 11, 2024, the Judicial Panel on Multidistrict Litigation granted a motion filed by Regeneron requesting transfer of Regeneron's patent infringement lawsuit pending against Amgen in the California Central District Court to the U.S. District Court for the Northern District of West Virginia (West Virginia District Court) for coordinated and consolidated pretrial proceedings with the five other cases involving EYLEA® biosimilars pending in that district.

On June 7, 2024, Regeneron filed a motion in the West Virginia District Court for a preliminary injunction to prohibit Amgen from engaging in the manufacture, use, offer for sell or sale within the United States, or importation into the United States, of PAVBLU until resolution of this lawsuit or the entry of a permanent injunction, whichever comes first. Regeneron's motion focused on U.S. Patent No. 11,084,865, a formulation patent. On September 23, 2024, the West Virginia District Court denied Regeneron's motion for a preliminary injunction, and Regeneron filed a notice of appeal, a motion to expedite the appeal, and an emergency motion for an injunction pending resolution of the appeal and for an administrative stay with the U.S. Court of Appeals for the Federal Circuit (Federal Circuit Court). On September 25, 2024, the Federal Circuit Court issued an order temporarily enjoining the launch of PAVBLU on an administrative basis while it considered Regeneron's motion for an injunction pending appeal. On October 22, 2024, the Federal Circuit Court denied Regeneron's motion for an injunction pending appeal and lifted the temporary injunction that was entered on September 25, 2024. Oral argument for the appeal was held on January 14, 2025. On March 14, 2025, the U.S. Court of Appeals for the Federal Circuit affirmed the denial by the West Virginia District Court of Regeneron's motion for a preliminary injunction.

On June 17, 2025, Regeneron filed a second lawsuit in the California Central District Court against Amgen alleging infringement of U.S. Patent No. 12,331,099 (the '099 Patent), also a formulation patent. By its complaint, Regeneron seeks, among other remedies, damages and an injunction prohibiting the commercial manufacture, use, offer for sale or sale in the United States or import into the United States of PAVBLU before the expiration of the '099 Patent. In July 2025, the Judicial Panel on Multidistrict Litigation issued an Order transferring the case from the California Central District Court to the West Virginia District Court for coordinated and consolidated pretrial proceedings with the other cases involving EYLEA® biosimilars pending in the district, and on July 31, 2025 the case was opened in the West Virginia District Court.

On September 12, 2025, Amgen responded to Regeneron's complaint asserting infringement of the '099 Patent, denying infringement and asserting counterclaims seeking a declaratory judgment that the '099 Patent is not infringed, invalid, and/or unenforceable, and counterclaims for Sherman Act (15 U.S.C. § 2) monopolization through Walker Process fraud, Sherman Act (15 U.S.C. § 2) attempted monopolization through Walker Process fraud, and unlawful and unfair practices under the California Unfair Competition Law. By its counterclaims, Amgen seeks, among other remedies, damages and an injunction against conduct by Regeneron. On September 29, 2025, the West Virginia District Court entered a scheduling order for the matters pending in the multidistrict litigation, including a claim construction hearing for November 23, 2026. On November 12, 2025, Regeneron filed a motion to strike certain of Amgen's affirmative defenses and to dismiss certain of Amgen's counterclaims, including the counterclaim seeking a declaratory judgment that the '099 Patent is unenforceable, the counterclaims for Sherman Act monopolization and attempted monopolization, and the counterclaim for unlawful and unfair practices under the California Competition Law.

On January 7, 2026, the West Virginia District Court entered a stipulation and order of dismissal that dismissed with prejudice Regeneron's claims with respect to 15 of the patents asserted in the 2024 Complaint and dismissed without prejudice Amgen's counterclaims and affirmative defenses with respect to those patents. Also on January 7, 2026, the West Virginia District Court granted Regeneron's motion for leave to amend the 2024 Complaint. The 2024 Complaint as amended alleges infringement of 19 patents and seeks, among other remedies, damages and an injunction prohibiting the commercial manufacture, use, offer for sale or sale in the United States or import into the United States of PAVBLU before the expiration of each of the patents found to be infringed.

KYPROLIS® (carfilzomib) Abbreviated New Drug Application (ANDA) Patent Litigation

Onyx Therapeutics, Inc. v. Amneal Pharmaceuticals of New York, LLC and Amneal EU, Limited.

On September 4, 2025, Onyx Therapeutics, Inc. (Onyx Therapeutics, a wholly-owned subsidiary of Amgen) filed a lawsuit in the U.S. District Court for the District of Delaware (Delaware District Court) against Amneal Pharmaceuticals of New York, LLC and Amneal EU, Limited (collectively, Amneal), asserting infringement of U.S. Patent No. 7,737,112 (the '112 Patent) based on Amneal's submission of an application pursuant to Section 505(b)(2) of the Federal Food, Drug, and Cosmetic Act seeking FDA approval to market a generic version of KYPROLIS. Onyx Therapeutics seeks an order from the

Delaware District Court making any FDA approval of Amneal's application effective no earlier than the expiration of the '112 Patent.

On November 6, 2025, Amneal answered the complaint and counterclaimed, seeking a declaration of non-infringement, and filed a motion for judgment on the pleadings of non-infringement. On December 8, 2025, Onyx answered Amneal's counterclaim, and on December 11, 2025, Onyx opposed Amneal's motion for judgment on the pleadings. On February 3, 2026, the Delaware District Court denied Amneal's motion for judgment on the pleadings.

TAVNEOS® (avacopan) Abbreviated New Drug Application (ANDA) Patent Litigation

ChemoCentryx, Inc. v. Zydus Pharmaceuticals (USA) Inc., Zydus Lifesciences Global FZE, and Zydus Lifesciences Limited

On January 16, 2026, ChemoCentryx, Inc. (ChemoCentryx, a wholly-owned subsidiary of Amgen) filed a lawsuit in the New Jersey District Court against Zydus Pharmaceuticals (USA) Inc., Zydus Lifesciences Global FZE, and Zydus Lifesciences Limited (collectively, Zydus) asserting infringement of U.S. Patent Nos. 11,603,356 (the '356 Patent) and 11,951,214 (the '214 Patent) based on Zydus's submission of an ANDA seeking FDA approval to market a generic version of TAVNEOS. ChemoCentryx seeks, among other things, an order from the New Jersey District Court making any FDA approval of Zydus's ANDA effective no earlier than the expiration of the '356 and '214 Patents.

ChemoCentryx, Inc. v. Annora Pharma Private Limited, Hetero USA Inc., and Hetero Labs Limited

On February 9, 2026, ChemoCentryx filed a lawsuit in the New Jersey District Court against Annora Pharma Private Limited, Hetero USA Inc., and Hetero Labs Limited (collectively, Annora) asserting infringement of the '356 Patent and the '214 Patent based on Annora's submission of an ANDA seeking FDA approval to market a generic version of TAVNEOS. ChemoCentryx seeks, among other things, an order from the New Jersey District Court making any FDA approval of Annora's ANDA effective no earlier than the expiration of the '356 and '214 Patents.

Antitrust Class Action

Regeneron Pharmaceuticals, Inc. Antitrust Action

On May 27, 2022, Regeneron filed suit against Amgen in the Delaware District Court for federal and state antitrust and unfair competition violations and tortious interference with prospective business relations. Regeneron alleges that Amgen's sales contracting practices for Repatha, ENBREL and Otezla with key insurers, third-party payers and PBMs have harmed the sales of its product PRALUENT® and focuses on two primary arguments: that Amgen improperly bundled sales of Repatha with ENBREL, Otezla and potentially other products and sought exclusive or de facto exclusive formulary positioning for Repatha. Amgen's initial responsive pleading, a motion to dismiss, was filed on August 1, 2022. The Delaware District Court denied Amgen's motion to dismiss the complaint.

On August 28, 2023, Regeneron filed its amended complaint, and on September 20, 2023, Amgen filed a counterclaim, alleging Regeneron's own anticompetitive conduct with respect to formulary position for Regeneron's drug, PRALUENT®, at CVS. Amgen notified the Delaware District Court that it would not be pursuing its counterclaim on April 24, 2025.

A jury trial was held in the Delaware District Court from May 5, 2025 to May 14, 2025. On May 15, 2025, the jury returned a verdict finding for Regeneron on its federal and state antitrust law and tortious interference claims but finding for Amgen on its below-cost pricing claim under California's Unfair Practices Act. The jury awarded Regeneron \$135.6 million in compensatory damages on its antitrust claims (which are subject to trebling under applicable law), or in the alternative, in compensatory damages plus \$271.2 million in punitive damages on its tortious interference claim, with such damages under either alternative claim totaling \$406.8 million. As Regeneron must elect between recovery under the antitrust or tortious interference claims, any potential damages award would be limited to one of these claims. Although we cannot predict with certainty the ultimate outcome of this litigation, Amgen believes that the jury's decision and amounts awarded are inconsistent with the law and evidence at trial.

Both parties have filed post-trial motions. On June 12, 2025, Amgen filed a renewed motion for judgment as a matter of law or, in the alternative, for a new trial. Also on June 12, 2025, Regeneron filed a motion for permanent injunctive relief, a constructive trust, and prejudgment interest. Both motions were fully briefed and a hearing on the post-trial motions was held on August 27, 2025.

In assessing whether we should accrue a liability for this litigation in our consolidated financial statements, we considered various factors, including the legal and factual circumstances of the case, the jury's award, the court's post-trial proceedings, applicable law, and the likelihood that the jury's award will be upheld after post-trial briefing and potentially on appeal. As a

result of this review, we have determined, in accordance with applicable accounting standards, that it is not probable that we will incur a loss as a result of this litigation, and we have therefore not recorded a liability for this matter.

The ultimate result of this litigation, however, is uncertain because it is reasonably possible that by settlement or final court judgment that none, some, or all of the jury's verdict and other relief sought might ultimately be awarded but the size of an award, if any, is not estimable at this time.

CareFirst of Maryland Antitrust Class Action

On August 6, 2024, CareFirst of Maryland, Inc., Group Hospitalization and Medical Services, Inc., and CareFirst BlueChoice, Inc. (collectively, CareFirst), filed a proposed class action lawsuit against Amgen Inc., Amgen Manufacturing, Limited (corrected to Amgen Manufacturing Limited LLC in CareFirst's amended complaint on filed October 11, 2024), and Immunex Corporation in the U.S. District Court for the Eastern District of Virginia (District Court for the Eastern District of Virginia), alleging federal and state antitrust claims and state consumer protection and unjust enrichment claims. CareFirst alleges that, in 2004, Amgen entered into an anticompetitive agreement with certain F. Hoffman-La Roche AG entities (Roche) and other parties that provided Amgen with rights to Roche's patents in a manner that enabled Amgen to allegedly unlawfully extend the life of patents applicable to ENBREL and, thereby, delay biosimilar entry. On November 4, 2024, Amgen filed a motion to dismiss, and CareFirst thereafter filed a second amended complaint on November 25, 2024. On January 8, 2025, Amgen filed a motion to dismiss the second amended complaint.

On September 30, 2025, the District Court for the Eastern District of Virginia issued an order granting in part and denying in part Amgen's motion to dismiss. The court dismissed CareFirst's antitrust claim under Puerto Rico law and its unjust enrichment claims under the laws of seven states and Puerto Rico, but otherwise permitted the claims to proceed. On October 20, 2025, Amgen filed a motion asking the district court to certify its order on the motion to dismiss for interlocutory appeal to the U.S. Court of Appeals for the Fourth Circuit. On November 3, 2025, CareFirst filed its opposition to Amgen's motion to certify for interlocutory appeal. On November 7, 2025, Amgen filed its answer to the second amended complaint, and on November 10, 2025, Amgen filed its reply in support of its motion to certify for interlocutory appeal.

Sandoz Inc. Antitrust Action

On April 11, 2025, Sandoz Inc. (Sandoz) filed a complaint in the U.S. District Court for the Eastern District of Virginia against Amgen Inc., Amgen Manufacturing Limited LLC, and Immunex Corporation claiming violations of the antitrust laws and tortious interference related to Amgen's patent rights to ENBREL, and seeking damages, injunctive relief and attorneys' fees. The factual allegations that form the basis for the claims of Sandoz's complaint are substantially similar to those asserted in the lawsuit filed against Amgen in the same court by CareFirst.

On June 20, 2025, Amgen filed a motion to dismiss the complaint. Sandoz filed its opposition to the motion to dismiss on July 21, 2025, and on August 21, 2025, Amgen filed its reply in support of the motion to dismiss.

Other Similar Antitrust Actions

In July and August 2025, seven lawsuits were filed in the California Superior Court in Ventura County, each alleging state law antitrust, consumer protection, and unjust enrichment claims, based on allegations substantially similar to those in the CareFirst class action. The cases were filed by: Centene Corporation, WellCare Health Plans, Inc., New York Quality Healthcare Corporation d/b/a Fidelis Care, and Health Net, LLC (collectively, Centene) on July 29, 2025; Humana Inc. (Humana) on July 29, 2025; Molina Healthcare, Inc. (Molina) on July 29, 2025; Blue Cross and Blue Shield of Florida, Inc. and Health Options, Inc. d/b/a Florida Blue HMO (collectively, BCBSFL) on July 29, 2025; Blue Cross and Blue Shield of Kansas City (BCBSKC) on July 29, 2025; Blue Cross and Blue Shield of Massachusetts, Inc. and Blue Cross and Blue Shield of Massachusetts HMO Blue, Inc. (collectively, BCBSMA) on August 8, 2025; and Health Care Services Corp. (HCSC) on August 21, 2025. Amgen subsequently removed the cases filed by BCBSFL, BCBSKC, BCBSMA, and HCSC to the California Central District Court. On September 29, 2025, HCSC voluntarily dismissed its case without prejudice. On October 1, 2025, BCBSFL, BCBSKC, and BCBSMA voluntarily dismissed their cases without prejudice.

On January 8, 2026, Amgen filed demurrers to the complaints filed by Centene, Humana, and Molina, seeking the dismissal of the claims in those complaints, with prejudice. On February 2, 2026, the complaint in the Centene case was amended to join BCBSFL, BCBSKC, BCBSMA, and HCSC as additional plaintiffs. Humana and Molina filed their opposition to Amgen's demurrers on February 5, 2026, and Amgen's replies are due on February 26, 2026. On February 5, Amgen filed a demurrer to the amended Centene complaint; Centene's opposition to the demurrer was filed February 9, 2026 and Amgen's reply is due February 26, 2026.

Amgen Inc. & Subsidiaries v. Commissioner of Internal Revenue

See Note 7, Income taxes, for discussion of the IRS tax dispute and the Company's petitions in the U.S. Tax Court.

Securities Class Action Litigation (Roofers Local No. 149 Pension Fund)

On March 13, 2023, Roofers Local No. 149 Pension Fund filed a purported class action (Roofers Securities Class Action) against Amgen, Robert Bradway and Peter Griffith in the U.S. District Court for the Southern District of New York (Southern District Court of New York). The action was brought on behalf of an alleged class of Amgen shareholders who owned stock between July 29, 2020 and April 27, 2022 (the alleged class period). Plaintiffs allege that the defendants made a series of materially false and misleading statements and omissions during the alleged class period regarding the failure to timely disclose the potential tax liability claimed by the IRS. Plaintiffs further allege that they and other purported class members suffered losses and damages resulting from declines in the market value of Amgen's common stock after the potential tax liability claimed by the IRS was disclosed.

On August 31, 2023, plaintiff filed an amended complaint and Amgen filed a motion to dismiss on November 6, 2023, which the Southern District Court of New York denied on September 30, 2024. On November 20, 2024, Amgen filed an answer to the amended complaint.

Pursuant to the Southern District Court of New York's order on September 11, 2025, the class certification briefing is due by April 24, 2026. The last day to file summary judgment motions is December 21, 2026, but no briefing schedule has been set and no trial date has been set.

On December 9, 2025, plaintiffs filed their motion for class certification. Amgen's opposition is due on March 9, 2026 and plaintiffs' reply is due May 8, 2026.

Shareholder Derivative Actions (Martin, Clearwater and DM Cohen)

On August 2, 2023, Leon Martin filed a derivative action (the Martin Derivative Action) captioned *Leon Martin v. Robert A. Bradway, et al.*, No. 1:23-cv-06754 (S.D.N.Y. Aug. 2, 2023), purportedly on behalf of Amgen, against Amgen, Robert Bradway, Peter Griffith and Amgen's independent Board members. The action was filed in the Southern District Court of New York as related to the pending Roofers Securities Class Action. The complaint in this matter alleges claims for violations of the Securities Exchange Act of 1934, breach of fiduciary duty, aiding and abetting breach of fiduciary duty, unjust enrichment and waste of corporate assets.

On December 7, 2023, Plaintiff filed a Notice of Voluntary Dismissal as to Board member Michael Drake.

On December 1, 2023, a second derivative action (the Clearwater Derivative Action) was filed, captioned *Cheri Clearwater v. Robert A. Bradway, et al.*, No. 1:23-cv-10538 (S.D.N.Y. Dec. 1, 2023), in the same court as the earlier-filed Martin Derivative Action. The second action is largely duplicative of the Martin Derivative Action, asserting the same claims purportedly on behalf of the Company against the individual directors that sat on Amgen's Board during the relevant time period (July 29, 2020 through April 27, 2022). The complaint asserts claims for breach of fiduciary duty, unjust enrichment, waste of corporate assets, abuse of control, gross mismanagement, and violations of Section 10(b) of the Exchange Act arising out of Amgen's disclosures with respect to its transfer pricing dispute with the IRS. However, the Clearwater Derivative Action complaint adds (1) two additional claims for violations of Sections 14(a) and 20(a) of the Exchange Act; (2) allegations that Amgen repurchased its own stock at artificially inflated prices during the relevant period; and (3) more detailed allegations as to why first making a demand on the Board would have been futile.

On January 16, 2024, the Southern District Court of New York consolidated the Martin Derivative Action and Clearwater Derivative Action (the Consolidated Action). The Southern District Court of New York entered an Order staying the Consolidated Action until a final judgment is entered in the Roofers Securities Class Action.

On February 12, 2025, DM Cohen, Inc. filed a third derivative action (the DM Cohen Derivative Action) captioned *DM Cohen, Inc. v. Robert A. Bradway, et al.*, No. 1:25-mc-00062 (S.D.N.Y. Feb. 12, 2025), purportedly on behalf of Amgen, against Amgen, Robert Bradway, Peter Griffith and Amgen's independent Board members. The DM Cohen Derivative Action was filed in the same court as the earlier filed Roofers Securities Class Action and the Consolidated Action. The complaint asserts claims for violations of the Securities Exchange Act of 1934, breach of fiduciary duty, waste of corporate assets and unjust enrichment.

The factual allegations that form the basis for the claims in the Consolidated Action and the DM Cohen Derivative Action are essentially the same as the allegations asserted in the Roofers Securities Class Action regarding purportedly false and

misleading statements and omissions made from July 29, 2020 through April 27, 2022 relating to Amgen's tax liabilities, business and finances, and the adequacy and maintenance of its internal controls.

On April 21, 2025, the Southern District Court of New York consolidated the derivative action filed by DM Cohen, Inc. with the Consolidated Action.

The case is stayed pending entry of a final judgment in the Roofers Securities Class action.

Shareholder Derivative Actions (Hamilton, Blackburn, Bryla)

On October 16, 2024, David Hamilton filed a derivative action in the Delaware Court of Chancery purportedly on behalf of Amgen, against nominal defendant Amgen, Robert Bradway, Peter Griffith and Amgen's Board members during the relevant time period (the Hamilton Derivative Action). The complaint in this matter alleges claims for breach of fiduciary duty and unjust enrichment.

On November 7, 2024, Charles Blackburn filed a derivative action in the Delaware Court of Chancery purportedly on behalf of Amgen, against nominal defendant Amgen, Robert Bradway, Peter Griffith and Amgen's Board members during the relevant time period (the Blackburn Derivative Action). The complaint alleges a claim for breach of fiduciary duty.

On December 6, 2024, Robert Bryla filed a derivative action in the Delaware Court of Chancery purportedly on behalf of Amgen, against nominal defendant Amgen, Robert Bradway, Peter Griffith and Amgen's Board members during the relevant time period (the Bryla Derivative Action). The complaint alleges claims for breach of fiduciary duty and unjust enrichment.

The factual allegations that form the basis for the claims in the Hamilton Derivative Action, Blackburn Derivative Action and Bryla Derivative Action are fundamentally the same as those asserted by the Roofers Local No. 149 Pension Fund on March 13, 2023 (alleging false and misleading statements and omissions made from July 29, 2020 through April 27, 2022 relating to Amgen's tax liabilities, business and finances, and the adequacy and maintenance of its internal controls).

On April 9, 2025, the Delaware Court of Chancery consolidated the derivative actions filed by each of David Hamilton, Charles Blackburn and Robert Bryla purportedly on behalf of Amgen against nominal defendant Amgen, Robert Bradway, Peter Griffith and Amgen's independent Board members.

The case is stayed pending entry of a final judgment in the Roofers Securities Class Action.

Shareholder Derivative Actions (Sieveking, Tierney)

On April 2, 2025, Carolyn Sieveking and James P. Tierney filed a derivative action (the Sieveking Derivative Action) in the Delaware District Court purportedly on behalf of Amgen, against nominal defendant Amgen, Robert Bradway, Peter Griffith, Linda Louie and Amgen's Board members during the relevant time period (the Sieveking Derivative Action). The complaint alleges claims for violations of Section 10(b), Rule 10b5 and Section 20(a) of the Securities Exchange Act of 1934, and breach of fiduciary duty.

The factual allegations that form the basis for the claims in the Sieveking Derivative Action is fundamentally the same as those asserted by the Roofers Local No. 149 Pension Fund.

On June 9, 2025, the Delaware District Court stayed the case pending entry of a final judgment in the Roofers Securities Class Action.

ChemoCentryx, Inc. Securities Matters

On May 5 and June 8 of 2021, ChemoCentryx and its Chief Executive Officer were named as defendants in two putative shareholder class actions filed in the U.S. District Court for the Northern District of California (Northern District Court of California). These cases were consolidated into *Homyk v. ChemoCentryx, Inc.* in which the plaintiffs allege violations of Sections 10(b) and 20(a) of the Securities Exchange Act in connection with statements regarding the New Drug Application for TAVNEOS and the underlying Phase 3 clinical trial, seeking an award of damages, interest and attorneys' fees. On March 28, 2022, the plaintiffs filed their consolidated amended complaint, and on May 19, 2022, ChemoCentryx moved to dismiss these claims.

On February 23, 2023, the Northern District Court of California substantially denied ChemoCentryx's motion to dismiss the matter in its entirety, while granting the motion to dismiss with respect to certain allegations of the plaintiffs. On August 25, 2023, the lead plaintiff moved to certify a class composed of all purchasers of ChemoCentryx stock between November 25, 2019 and May 6, 2021.

On March 6, 2024, the Northern District Court of California certified a class of all persons who purchased or otherwise acquired the common stock of ChemoCentryx between November 26, 2019 and May 6, 2021. The deadline for class members to opt out of the class action was January 14, 2025.

On August 15, 2025, the Northern District Court of California granted defendants' motion for summary judgment in its entirety and denied lead plaintiff's motion for partial summary judgment. On September 12, 2025, the lead plaintiff filed a notice of appeal to the Ninth Circuit Court of Appeals, and filed its opening brief on January 5, 2026.

On January 27, 2026, the parties informed the Court of Appeals they had reached an agreement to settle the case on a class-wide basis, subject to approval by the district court.

Opt-Out Cases

Prior to the opt-out deadline, on May 2, 2024, RA Capital Healthcare Fund, LP filed two securities cases (which are similar to the class action), in the California Superior Court in Ventura County and in the Northern District Court of California, against ChemoCentryx and its former Chief Executive Officer, Dr. Thomas Schall. On July 2, 2024, the state court stayed the case pending an order on summary judgment in the federal class action. Defendants, including ChemoCentryx, moved to dismiss the complaint, and on June 13, 2025, the court issued an order staying the federal case pending resolution of the class action. The court did not reach the merits of defendants' motion to dismiss.

Commitments – U.S. repatriation tax

Under the 2017 Tax Act, we elected to pay in eight annual installments the repatriation tax related primarily to prior indefinitely invested earnings of our foreign operations. The final U.S. repatriation tax payment of \$1.8 billion was made in 2025.

AMGEN INC.

VALUATION AND QUALIFYING ACCOUNTS

Years ended December 31, 2025, 2024 and 2023

(In millions)

Allowance for doubtful accounts	Balance at beginning of period	Additions charged to costs and expenses	Deductions	Other changes	Balance at end of period
2025	\$ 38	\$ 1	\$ —	\$ 4	\$ 43
2024	\$ 28	\$ 13	\$ —	\$ (3)	\$ 38
2023	\$ 22	\$ 6	\$ —	\$ —	\$ 28

**DESCRIPTION OF AMGEN INC.'S SECURITIES
REGISTERED PURSUANT TO SECTION 12 OF THE
SECURITIES EXCHANGE ACT OF 1934**

As of February 10, 2026, Amgen Inc. has two classes of securities registered under Section 12 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"): (1) our common stock, par value \$0.0001 per share (the "Common Stock"); and (2) our 2.000% Senior Notes due 2026 (the "Notes").

DESCRIPTION OF COMMON STOCK

The following description of our capital stock is a summary and does not purport to be complete. It is subject to and qualified in its entirety by reference to our restated certificate of incorporation, as amended ("certificate of incorporation") and our amended and restated bylaws, each of which are incorporated by reference as an exhibit to the Annual Report on Form 10-K ("Annual Report"). The terms "Amgen" "we," "our," and "us" refer solely to Amgen Inc. and not its subsidiaries.

Our authorized capital stock includes 2,750,000,000 shares of Common Stock. Each holder of our Common Stock is entitled to one vote per share on all matters to be voted upon by our stockholders. Upon any liquidation, dissolution or winding up of our business, the holders of our Common Stock are entitled to share equally in all assets available for distribution after payment of all liabilities, subject to the liquidation preference of shares of preferred stock, if any, then outstanding. Our Common Stock has no preemptive or conversion rights. All outstanding shares of common stock are fully paid and non-assessable. Our outstanding shares of common stock are quoted on the Nasdaq Global Select Market under the symbol "AMGN."

Dividends

Subject to preferences that may be applicable to any preferred stock (if any such stock be issued and outstanding), the holders of Common Stock are entitled ratably to receive dividends, if any, declared by our board of directors out of funds legally available for the payment of dividends.

Anti-Takeover Effects of Delaware Law

We are subject to the provisions of Section 203 of the Delaware General Corporation Law. Under Section 203, we would generally be prohibited from engaging in any business combination with any interested stockholder for a period of three years following the time that this stockholder became an interested stockholder unless:

- prior to such time, the board of directors of the corporation approved either the business combination or the transaction that resulted in the stockholder becoming an interested stockholder;
- upon consummation of the transaction which resulted in the stockholder becoming an interested stockholder, the interested stockholder owned at least 85% of the voting stock of the corporation outstanding at the time the transaction commenced, excluding for purposes of determining the voting stock outstanding (but not the outstanding voting stock owned by the interested stockholder) those shares owned (i) by persons who are directors and also officers and (ii) employee stock plans in which employee participants do not have the right to determine confidentially whether shares held subject to the plan will be tendered in a tender or exchange offer; or
- at or subsequent to such time, the business combination is approved by the board of directors and authorized at an annual or special meeting of stockholders, and not by written consent, by the affirmative vote of at least 66 2/3% of the outstanding voting stock which is not owned by the interested stockholder.

Under Section 203, a "business combination" includes:

- any merger or consolidation involving the corporation and the interested stockholder;
- any sale, lease, exchange, mortgage, pledge, transfer or other disposition of 10% or more of the assets of the corporation involving the interested stockholder;

- any transaction which results in the issuance or transfer by the corporation or by any direct or indirect majority-owned subsidiary of the corporation of any stock of the corporation or of such subsidiary to the interested stockholder, subject to limited exceptions;
- any transaction involving the corporation or any direct or indirect majority-owned subsidiary of the corporation which has the effect, directly or indirectly, of increasing the proportionate share of the stock of any class or series, or securities convertible into the stock of any class or series, of the corporation or of any such subsidiary which is owned by the interested stockholder; or
- the receipt by the interested stockholder of the benefit of any loans, advances, guarantees, pledges or other financial benefits provided by or through the corporation or any direct or indirect majority-owned subsidiary of the corporation.

In general, Section 203 defines an interested stockholder as an entity or person beneficially owning 15% or more of the outstanding voting stock of the corporation and any entity or person affiliated with or controlling or controlled by such entity or person.

Transfer Agent

The transfer agent and registrar for our Common Stock is the American Stock Transfer & Trust Company.

DESCRIPTION OF THE NOTES

The following description of our Notes is a summary and does not purport to be complete. It is subject to and qualified in its entirety by reference to the indenture, dated as of May 22, 2014 (the “Indenture”), between us and The Bank of New York Mellon Trust Company, N.A., as trustee (the “Trustee”), which are incorporated by reference as exhibits to the Annual Report of which this Exhibit 4.32 is a part. The Notes are traded on The Nasdaq Stock Market LLC under the trading symbol of “AMGN26.” We encourage you to read the above referenced Indenture for additional information.

General

We issued €750,000,000 in aggregate principal amount of 2.000% Senior Notes, maturing February 25, 2026 and bearing interest at a rate of 2.000% per annum, payable annually on February 25 of each year. As of February 10, 2026, €750,000,000 aggregate principal amount of the Notes was outstanding.

We may, without notice to or the consent of the holders or beneficial owners of the Notes of any series, create and issue additional Notes and/or notes having the same ranking, interest rate, maturity and other terms as the Notes of that series. Any additional debt securities having such similar terms, together with that series of Notes, could be considered part of the same series of Notes under the Indenture; *provided* that, in the case of any notes represented by global notes, for so long as may be required by the United States Securities Act of 1933, as amended (the “Securities Act”), or the procedures of the common depositary, the Euroclear System (“Euroclear”) or Clearstream Banking, S.A. (“Clearstream”) (or a successor or clearing system), such additional Notes will be represented by one or more separate global notes in accordance with the terms of the Indenture and subject to applicable transfer or other restrictions.

The Notes are redeemable prior to maturity as described below under the headings “—Optional Redemption” and “—Redemption Upon Changes in Withholding Taxes.” The Notes do not have the benefit of any sinking funds. The Notes of each series are issued only in registered form without coupons attached in minimum denominations of €100,000 and any integral multiple of €1,000 in excess thereof. Each series of Notes are represented by one or more global securities deposited with, or on behalf of, a common depositary for Euroclear and Clearstream (the “global notes”).

Certain Definitions

As used herein, the following terms have the meanings set forth below.

“*Attributable Liens*” means in connection with a sale and lease-back transaction the lesser of:

- (1) the fair market value of the assets subject to such transaction; and
- (2) the present value (discounted at a rate per annum equal to the average interest borne by all outstanding debt securities issued under the Indenture (which may include debt securities in addition to the Notes) determined on a weighted average basis and compounded semi-annually) of the obligations of the lessee for rental payments during the term of the related lease.

“*Business Day*” means any day on which commercial banks and foreign exchange markets are open for business in New York and London and which is a day on which the Trans-European Automated Real-Time Gross Settlement Express Transfer System (TARGET2) is operating.

“*Calculation Agent*” means an independent financial institution appointed by Amgen, which may include the paying agent, any of the managers or their respective affiliates who agree to serve in such capacity.

“*Capital Lease*” means any Indebtedness represented by a lease obligation of a Person incurred with respect to real property or equipment acquired or leased by such Person and used in its business that is required to be recorded as a capital lease in accordance with GAAP.

“*Consolidated Net Worth*” means, as of any date of determination, the Stockholders’ Equity of us and our Consolidated Subsidiaries on that date.

“*Consolidated Subsidiary*” means, as of any date of determination and with respect to any Person, any Subsidiary of that Person whose financial data is, in accordance with GAAP, reflected in that Person’s consolidated financial statements.

“*Credit Facilities*” means, one or more debt facilities (including, without limitation, the revolving credit agreement and the term loan credit agreement, as applicable) or commercial paper facilities, in each case, with banks or other institutional lenders providing for revolving credit loans, term loans, receivables financing (including through the sale of receivables to such lenders or to special purpose entities formed to borrow from such lenders against such receivables) or letters of credit, in each case, as amended, restated, modified, renewed, refunded, replaced (whether upon or after termination or otherwise) or refinanced (including by means of sales of debt securities to institutional investors) in whole or in part from time to time.

“*Exempted Debt*” means the sum of the following as of the date of determination:

- (1) our Indebtedness incurred after the first issue date of the Notes and secured by Liens not permitted by the first sentence under “—Limitation on Liens” below; and
- (2) our and our Subsidiaries’ Attributable Liens in respect of sale and lease-back transactions entered into after the first issue date of the Notes pursuant to the second paragraph of “—Limitation on Sale and Lease-Back Transactions” below.

“*GAAP*” means accounting principles generally accepted in the United States set forth in the Accounting Standards Codification of the Financial Accounting Standards Board or in such other documents by such other entity as have been approved by a significant segment of the accounting profession, which are in effect as of the date of determination.

“*Governmental Agency*” means:

- (1) any foreign, federal, state, county or municipal government, or political subdivision thereof;
- (2) any governmental or quasi-governmental agency, authority, board, bureau, commission, department, instrumentality or public body;
- (3) any court or administrative tribunal; and
- (4) with respect to any Person, any arbitration tribunal or other nongovernmental authority to whose jurisdiction that Person has consented.

“*Hedging Obligations*” means, with respect to any specified Person, the obligations of such Person under:

- (1) interest rate swap agreements (whether from fixed to floating or from floating to fixed), interest rate cap agreements and interest rate collar agreements;
- (2) other agreements or arrangements designed to manage interest rates or interest rate risk; and
- (3) other agreements or arrangements designed to protect such Person against fluctuations in currency exchange rates or commodity prices.

“*Indebtedness*” of any Person means, without duplication, any indebtedness, whether or not contingent, in respect of borrowed money or evidenced by bonds, notes, debentures or similar instruments or letters of credit (or reimbursement agreements with respect thereto) or representing the balance deferred and unpaid of the purchase price of any Property (including pursuant to Capital Leases), except any such balance that constitutes an accrued expense or trade payable, if and to the extent any of the foregoing indebtedness would appear as a liability upon a balance sheet of such Person prepared on a consolidated basis in accordance with GAAP (but does not include

contingent liabilities which appear only in a footnote to a balance sheet), and shall also include, to the extent not otherwise included, the guaranty of items which would be included within this definition.

“*Laws*” means, collectively, all foreign, federal, state and local statutes, treaties, rules, regulations, ordinances, codes and administrative or controlling precedents of any Governmental Agency.

“*Lien*” means any lien, security interest, charge or encumbrance of any kind (including any conditional sale or other title retention agreement, any lease in the nature thereof, and any agreement to give any security interest).

“*Make-Whole Amount*” means the excess of (1) the net present value, on the redemption date, of the principal being redeemed or paid and the amount of interest (exclusive of interest accrued to the date of redemption) that would have been payable if such redemption had not been made, over (2) the aggregate principal amount of the Notes being redeemed or paid. Net present value shall be determined by discounting, on a semi-annual basis, such principal and interest at the Reinvestment Rate (as defined below and as determined on the third Business Day preceding the date such notice of redemption is given) from the respective dates on which such principal and interest would have been payable if such redemption had not been made.

“*Permitted Liens*” means:

- (1) Liens securing Indebtedness under Credit Facilities;
- (2) Liens on accounts receivable, merchandise inventory, equipment, and patents, trademarks, trade names and other intangibles, securing our Indebtedness;
- (3) Liens on any of our assets, any of our Subsidiaries’ assets, or the assets of any joint venture to which we or any of our Subsidiaries is a party, created solely to secure obligations incurred to finance the refurbishment, improvement or construction of such asset, which obligations are incurred no later than 24 months after completion of such refurbishment, improvement or construction, and all renewals, extensions, refinancings, replacements or refundings of such obligations;
- (4) (a) Liens given to secure the payment of the purchase price incurred in connection with the acquisition (including acquisition through merger or consolidation) of Property (including shares of stock), including Capital Lease transactions in connection with any such acquisition, and
(b) Liens existing on Property at the time of acquisition thereof or at the time of acquisition by us or one of our Subsidiaries of any Person then owning such Property whether or not such existing Liens were given to secure the payment of the purchase price of the Property to which they attach; provided that, with respect to clause (a), the Liens shall be given within 24 months after such acquisition and shall attach solely to the Property acquired or purchased and any improvements then or thereafter placed thereon;
- (5) Liens in favor of customs and revenue authorities arising as a matter of law to secure payment of customs duties in connection with the importation of goods;
- (6) Liens upon specific items of inventory or other goods and proceeds of any Person securing such Person’s obligations in respect of bankers’ acceptances issued or created for the account of such Person to facilitate the purchase, shipment or storage of such inventory or other goods;
- (7) Liens securing reimbursement obligations with respect to letters of credit that encumber documents and other Property relating to such letters of credit and the products and proceeds thereof;

- (8) Liens on key-man life insurance policies granted to secure our Indebtedness against the cash surrender value thereof;
- (9) Liens encumbering customary initial deposits and margin deposits and other Liens in the ordinary course of business, in each case securing Hedging Obligations and forward contract, option, futures contracts, futures options or similar agreements or arrangements designed to protect us or any of our Subsidiaries from fluctuations in interest rates, currencies or the price of commodities;
- (10) Liens arising out of conditional sale, title retention, consignment or similar arrangements for the sale of goods entered into by us or any of our Subsidiaries in the ordinary course of business;
- (11) pre-existing Liens on assets acquired by us or any of our Subsidiaries after the first issue date of the Notes;
- (12) Liens in our favor or the favor of any of our Subsidiaries;
- (13) inchoate Liens incident to construction or maintenance of real property, or Liens incident to construction or maintenance of real property, now or hereafter filed of record for sums not yet delinquent or being contested in good faith, if reserves or other appropriate provisions, if any, as shall be required by GAAP shall have been made therefor;
- (14) statutory Liens arising in the ordinary course of business with respect to obligations which are not delinquent or are being contested in good faith, if reserves or other appropriate provisions, if any, as shall be required by GAAP shall have been made therefor;
- (15) Liens consisting of pledges or deposits to secure obligations under workers' compensation laws or similar legislation, including Liens of judgments thereunder which are not currently dischargeable;
- (16) Liens consisting of pledges or deposits of Property to secure performance in connection with operating leases made in the ordinary course of business to which we or any of our Subsidiaries is a party as lessee, provided the aggregate value of all such pledges and deposits in connection with any such lease does not at any time exceed 16 2/3% of the annual fixed rentals payable under such lease;
- (17) Liens consisting of deposits of Property to secure our statutory obligations or statutory obligations of any of our Subsidiaries in the ordinary course of its business;
- (18) Liens consisting of deposits of Property to secure (or in lieu of) surety, appeal or customs bonds in proceedings to which we or any of our Subsidiaries is a party in the ordinary course of its business, but not in excess of \$75,000,000;
- (19) purchase money Liens or purchase money security interests upon or in any Property acquired or held by us or any of our Subsidiaries in the ordinary course of business to secure the purchase price of such Property or to secure indebtedness incurred solely for the purpose of financing the acquisition of such Property;

(20) Liens on an asset created in connection with the acquisition, construction or development of additions, extensions or improvements to such asset which shall be financed by obligations described in Sections 142, 144(a) or 144(c) of the Code, or by obligations entitled to substantially similar tax benefits under other legislation or regulations in effect from time to time; and

(21) Liens on Property subject to escrow or similar arrangements established in connection with litigation settlements.

“*Person*” means any individual, corporation, partnership, joint venture, association, limited liability company, joint-stock company, trust, unincorporated organization or government or any agency or political subdivision thereof.

“*Property*” means any property or asset, whether real, personal or mixed, or tangible or intangible.

“*Reference Bund*” means the Federal Government Bond of Bundesrepublik Deutschland due February 15, 2026, with ISIN 0001102390.

“*Reference Dealers*” means each of the four banks selected by a Calculation Agent which are primary European government security dealers, and their respective successors, or market makers in pricing corporate bond issues.

“*Reinvestment Rate*” means 0.300% plus the average of the four quotations given by the Reference Dealers of the mid-market annual yield to maturity of the Reference Bund at 11: 00 a.m. (Central European time (“CET”)) on the fourth Business Day preceding such redemption date and if the Reference Bund is no longer outstanding, a Similar Security will be chosen by the Calculation Agent at 11: 00 a.m. (CET) on the third Business Day in London preceding such redemption date, quoted in writing by the Calculation Agent to us.

“*Similar Security*” means a reference bond or reference bonds issued by the German Federal Government having an actual or interpolated maturity comparable with the remaining term of the Notes that would be utilized, at the time of selection and in accordance with customary financial practice, in pricing new issues of corporate debt securities of comparable maturity to the remaining term of the Notes.

“*Stockholders’ Equity*” means, as of any date of determination, stockholders’ equity as of that date determined in accordance with GAAP; provided that there shall be excluded from Stockholders’ Equity any amount attributable to capital stock that is, directly or indirectly, required to be redeemed or repurchased by the issuer thereof at a specified date or upon the occurrence of specified events or at the election of the holder thereof.

“*Subsidiary*” of any specified person means any corporation, association or other business entity of which more than 50% of the total voting power of shares of capital stock entitled (without regard to the occurrence of any contingency) to vote in the election of directors, managers or trustees thereof is at the time owned or controlled, directly or indirectly, by such person or one or more of the other Subsidiaries of that person or a combination thereof.

Paying Agent and Registrar

The Bank of New York Mellon, London Branch, is the principal paying agent for the Notes (the “principal paying agent”). The Bank of New York Mellon Trust Company, N.A., is the security registrar for the Notes. Upon notice to the Trustee, we may change any paying agent or security registrar, and we or any of our subsidiaries may act as paying agent or registrar.

Interest

The Notes accrue interest at a rate of 2.000% per annum. The Notes accrue interest on their stated principal amounts from the most recent interest payment date on which interest has been paid or duly provided for. Accrued and unpaid interest on the Notes are payable annually in arrears on February 25 of each year. In each case, interest is paid to the holder in whose name a note is registered at the close of business on the day that is one Business Day prior to the relevant interest payment date.

Interest on the Notes is computed on the basis of the actual number of days in the period for which interest is being calculated and the actual number of days from and including the last date on which interest was paid on the Notes, to but excluding the next scheduled interest payment date. This payment convention is referred to as Actual/

Actual (ICMA) as defined in the rulebook of the International Capital Market Association. If any date on which interest, principal or premium is payable on the Notes is not a Business Day, then payment of such amounts payable on such date will be made on the next succeeding day that is a Business Day (and, except as provided under “—Payment of Additional Amounts,” without any interest or other payment in respect of any such delay) with the same force and effect as if made on such interest payment date or maturity date, as the case may be.

Any amounts payable on any Notes that are not punctually paid on any payment date will cease to be payable to the person in whose name such Notes are registered on the relevant record date, and such defaulted payment will instead be payable to the person in whose name such Notes are registered on the special record date or other specified date determined in accordance with the Indenture.

Ranking

The Notes are senior unsecured obligations of Amgen. The Notes rank:

- equal in right of payment to all of our other existing and future senior unsecured indebtedness;
- senior in right of payment to all of our existing and future subordinated indebtedness; and
- effectively subordinated in right of payment to all of our subsidiaries’ obligations (including secured and unsecured obligations) and subordinated in right of payment to our secured obligations, to the extent of the assets securing such obligations.

The Notes and the Indenture do not limit our ability to incur additional indebtedness. We may incur substantial additional amounts of indebtedness in the future.

Optional Redemption

The Notes may be redeemed prior to maturity at our option, at any time in whole or from time to time in part. If the Notes are redeemed before November 25, 2025 (three months prior to the maturity date of the Notes), the redemption price will equal the sum of (1) 100% of the principal amount being redeemed, plus accrued and unpaid interest to, but not including, the redemption date, and (2) the Make-Whole Amount, if any. If the Notes are redeemed on or after November 25, 2025 (three months prior to the maturity date of the Notes), the redemption price will equal 100% of the principal amount being redeemed, plus accrued and unpaid interest to, but not including, the redemption date.

If we give notice as provided in the Indenture and funds for the redemption of any Notes called for redemption sufficient to pay the redemption price have been deposited with the principal paying agent on or before 10:00 a.m., London time, on the redemption date, such Notes will cease to bear interest on the date fixed for redemption. Thereafter, the only right of the holders of such Notes will be to receive payment of the redemption price.

Upon surrender of a note that is redeemed in part, we shall execute and the Trustee shall authenticate for the holder a new note of the same series and the same maturity equal in principal amount to the unredeemed portion of the note surrendered.

The Notes are redeemable prior to maturity as described below under the headings “—Optional Redemption” and “—Redemption Upon Changes in Withholding Taxes.” The Notes do not have the benefit of any sinking funds. The Notes of each series are issued only in registered form without coupons attached in minimum denominations of €100,000 and any integral multiple of €1,000 in excess thereof. Each series of Notes are represented by one or more global securities deposited with, or on behalf of, a common depositary for Euroclear and Clearstream (the “global notes”).

Payments on the global notes are made through the principal paying agent (as defined herein under the heading “—Paying Agent and Registrar”). Payments on the Notes are made at the specified office or agency of the principal paying agent; *provided* that all such payments with respect to Notes represented by one or more global notes registered in the name of or held by a nominee of Euroclear or Clearstream, as applicable, will be by wire transfer of immediately available funds to the account specified by the holder or holders thereof.

In addition, at our option, if certificated notes are issued, we may make payments by check mailed to the holder’s registered address or by wire transfer to the account shown on the register for the certificated notes.

If certificated notes are issued, they will be issued only in minimum denominations of €100,000 principal amount and integral multiples of €1,000 in excess thereof upon receipt by the applicable registrar of instructions relating thereto and any certificates and other documentation required under the Indenture. It is expected that such instructions will be based upon directions received by Euroclear or Clearstream, as applicable, from the participant which owns the relevant book-entry interests. Certificated notes issued in exchange for book-entry interests will, except as provided in the Indenture, be subject to, and will have a legend with respect to the restrictions on transfer summarized below.

Subject to the restrictions on transfer referred to above, Notes issued as certificated notes may be transferred or exchanged, in whole or in part, in minimum denominations of €100,000 principal amount and integral multiples of €1,000 in excess thereof to persons who take delivery thereof in the form of certificated notes. In connection with any such transfer or exchange, the Indenture requires the transferring or exchanging holder to, among other things, furnish appropriate endorsements and transfer documents, to furnish information regarding the account of the transferee at Euroclear or Clearstream, where appropriate, to furnish certain certificates and opinions, and to pay any tax or other governmental charge in connection with such transfer or exchange. Any such transfer or exchange will otherwise be made without charge to the holder.

Notwithstanding the foregoing, we are not required to register the transfer or exchange of any Notes:

- for a period of 15 days prior to any date fixed for the redemption of the Notes;
- for a period of 15 days immediately prior to the date fixed for selection of Notes to be redeemed in part;
- for a period of 15 days prior to the record date with respect to any interest payment date; or
- which the holder has tendered (and not withdrawn) for repurchase in connection with a change of control offer.

Redemption Upon Changes in Withholding Taxes

If (a) as a result of any change in, or amendment to, the laws (or any regulations or rulings promulgated thereunder) of the United States (or any political subdivision or taxing authority thereof or therein having power to tax) (a “Relevant Taxing Jurisdiction”), or any change in, or amendment to, the official position regarding the application or interpretation of such laws, regulations or rulings (including by virtue of a holding, judgment or order by a court of competent jurisdiction or a change in published administrative practice), which change or amendment is announced on or after the date of the applicable prospectus supplement, we become or will become obligated to pay additional amounts as described herein under the heading “—Payment of Additional Amounts” or (b) any act is taken by a Relevant Taxing Jurisdiction on or after the date of the applicable prospectus supplement, whether or not such act is taken with respect to us or any affiliate, that results in a substantial probability that we will or may be required to pay such additional amounts, then we may, at our option, redeem the Notes of any affected series, as a whole but not in part, upon not less than 15 days’ nor more than 60 days’ published notice in accordance with the applicable notice requirement, at 100% of their principal amount, together with interest accrued thereon to the date fixed for redemption; *provided* that we determine, in our business judgment, that the obligation to pay such additional amounts cannot be avoided by the use of reasonable measures available to us (which does not include substitution of the obligor under the Notes). No redemption pursuant to (a) or (b) above may be made unless we have received an opinion of independent counsel to the effect that as a result of such change or amendment we will, or that an act taken by a Relevant Taxing Jurisdiction has resulted in a substantial probability that we will, or may, be required to pay the additional amounts described herein under the heading “—Payment of Additional Amounts,” and we shall have delivered to the Trustee a certificate, signed by a duly authorized officer, stating that based on such opinion we are entitled to redeem the Notes pursuant to their terms.

Notice of Redemption

We will publish a notice of any redemption of any affected series of Notes described above in accordance with the applicable notice provisions. If fewer than all of the Notes are to be redeemed at any time, the principal paying agent will select the Notes to be redeemed in accordance with the rules of the principal securities exchange, if any, on which the Notes are listed at such time or, if the not Notes are not listed on a securities exchange, in accordance with the rules of Euroclear or Clearstream, or absent any such rules, *pro rata*, by lot; *provided, however*, that no such partial redemption shall reduce the portion of the principal amount of a note not redeemed to less than

€100,000. The principal paying agent shall not be liable for any selections made by it in accordance with this paragraph.

We will give notice of any optional redemption to the registered holders of Notes at least 15 but not more than 60 days before a redemption date. The notice shall identify the Notes to be redeemed and shall state:

- the redemption date;
- the redemption price;
- the name and address of the paying agent;
- if any Notes are being redeemed in part, the portion of the principal amount of such notes to be redeemed and that, after the redemption date and upon surrender of such Notes, a new note or notes in principal amount equal to the unredeemed portion of the original note shall be issued in the name of the holder of the Notes thereof upon cancellation of the original note;
- that the notes called for redemption must be surrendered to the paying agent to collect the redemption price;
- that interest on the Notes called for redemption ceases to accrue on and after the redemption date unless we default in the deposit of the redemption price; and
- the CUSIP and/or ISIN number of the Notes.

At our request, the Trustee shall give the notice of redemption in our name and at our expense.

Payment of Additional Amounts

All payments of principal and interest on the Notes will be made free and clear of and without withholding or deduction for or on account of any present or future tax, assessment or other governmental charge (collectively, “Taxes”) imposed by any Relevant Taxing Jurisdiction, unless the withholding of such Taxes is required by law or the official interpretation or administration thereof. We will, subject to the exceptions and limitations set forth below, pay such additional amounts as are necessary in order that the net payment of the principal of and interest on the applicable series of Notes to a holder who is not a U.S. person for U.S. federal income tax purposes, after deduction for any present or future Taxes of any Relevant Taxing Jurisdiction, imposed by withholding with respect to the payment, will not be less than the amount provided in such Notes to be then due and payable; *provided, however*, that the foregoing obligation to pay additional amounts shall not apply:

(1) to any Taxes that are imposed or withheld solely by reason of the holder or beneficial owner, or a fiduciary, settlor, beneficiary, member or shareholder of the holder if the holder is an estate, trust, partnership or corporation, or a person holding a power over an estate or trust administered by a fiduciary holder, being considered as:

- (a) being or having been present or engaged in a trade or business in the United States or having or having had a permanent establishment in the United States;
- (b) having a current or former relationship with the United States, including a relationship as a citizen or resident thereof;
- (c) being or having been a foreign or domestic personal holding company, a passive foreign investment company or a controlled foreign corporation with respect to the United States or a corporation that has accumulated earnings to avoid U.S. federal income tax;
- (d) being or having been a “10-percent shareholder” of the obligor under the Notes within the meaning of section 871(h)(3) of the U.S. Internal Revenue Code of 1986, as amended (the “Code”), or any successor provision; or
- (e) being or having been a bank receiving interest described in section 881(c)(3)(A) of the Code or any successor provision;

(2) to any holder that is not the sole beneficial owner of the note, or a portion thereof, or that is a fiduciary or partnership, but only to the extent that a beneficiary or settlor with respect to the fiduciary, a beneficial owner or member of the partnership would not have been entitled to the payment of an additional amount had the beneficiary, settlor, beneficial owner or member received directly its beneficial or distributive share of the payment;

(3) to any Taxes that are imposed or withheld solely by reason of the failure to (a) comply with certification, identification or information reporting requirements concerning the nationality, residence, identity or connection with a Relevant Taxing Jurisdiction of the holder or beneficial owner of such note, if compliance is required by statute or by regulation of the Relevant Taxing Jurisdiction as a precondition to relief or exemption from such Taxes (including the submission of an applicable U.S. Internal Revenue Service (“IRS”) Form W-8 (with any required attachments)) or (b) comply with any informational gathering and reporting requirements or to take any similar action (including entering into any agreement with the IRS), in each case, that are required to obtain the maximum available exemption from withholding by a Relevant Taxing Jurisdiction that is available to payments received by or on behalf of the holder;

(4) to any Taxes that are imposed otherwise than by withholding from the payment;

(5) to any Taxes that are imposed or withheld solely by reason of a change in law, regulation, or administrative or judicial interpretation that becomes effective more than 15 days after the payment becomes due or is duly provided for, whichever occurs later;

(6) to any estate, inheritance, gift, sales, excise, transfer, wealth or personal property tax or a similar tax, assessment or governmental charge;

(7) to any Taxes required to be withheld by any paying agent from any payment of principal of or interest on any note, if such payment can be made without such withholding by any other paying agent;

(8) to any Taxes that are imposed or levied by reason of the presentation (where presentation is required in order to receive payment) of such notes for payment on a date more than 30 days after the date on which such payment became due and payable, except to the extent that the holder or beneficial owner thereof would have been entitled to additional amounts had the notes been presented for payment on any date during such 30 day period;

(9) to any Taxes that are imposed or withheld pursuant to Sections 1471 through 1474 of the Code, as of the issue date (or any amended or successor version of such sections), any U.S. Treasury Regulations promulgated thereunder, any official interpretations thereof, any similar law or regulation adopted pursuant to an intergovernmental agreement between a non-U.S. jurisdiction and the United States with respect to the foregoing or any agreements entered into pursuant to Section 1471(b)(1) of the Code; or

(10) in the case of any combination of any items (1) through (9).

The notes are subject in all cases to any tax, fiscal or other law or regulation or administrative or judicial interpretation applicable thereto. Except as specifically provided under this heading “—Payment of Additional Amounts,” we are not required to make any payment with respect to any tax, assessment or governmental charge imposed by any government or a political subdivision or taxing authority thereof or therein.

Change of Control Offer

If a change of control triggering event occurs, unless we have exercised our option to redeem the notes as described above, we will be required to make an offer (the “change of control offer”) to each holder of the notes to repurchase all or any part (equal to €100,000 or integral multiples of €1,000 in excess thereof) of that holder’s notes on the terms set forth in such notes. In the change of control offer, we will be required to offer payment in cash equal to 101 % of the aggregate principal amount of notes repurchased, plus accrued and unpaid interest, if any, on the notes repurchased to the date of repurchase (the “change of control payment”). Within 30 days following any change of control triggering event, a notice will be provided to holders of the notes describing the transaction that constitutes the change of control triggering event and offering to repurchase the notes on the date specified in the notice, which date will be no earlier than 30 days and no later than 60 days from the date such notice is provided (the “change of control payment date”); provided, however, that in no event will the change of control payment date occur prior to the date 90 days following the first issue date of the notes.

On the change of control payment date, we will, to the extent lawful:

- accept for payment all notes or portions of notes properly tendered pursuant to the change of control offer;
- by 10:00 a.m., London time, deposit with the principal paying agent an amount equal to the change of control payment in respect of all notes or portions of notes properly tendered; and
- deliver or cause to be delivered to the Trustee the notes properly accepted together with an officer’s certificate stating the aggregate principal amount of notes or portions of notes being repurchased.

We will not repurchase any notes if there has occurred and is continuing on the change of control payment date an event of default under the Indenture, other than a default in the payment of the change of control payment upon a change of control triggering event.

We will comply with the requirements of Rule 14e-1 under the U.S. Securities Exchange Act of 1934, as amended (the “Exchange Act”), and any other securities laws and regulations thereunder to the extent those laws and regulations are applicable in connection with the repurchase of the notes as a result of a change of control triggering event. To the extent that the provisions of any such securities laws or regulations conflict with the change of control offer provisions of the notes, we will comply with those securities laws and regulations and will not be deemed to have breached our obligations under the change of control offer provisions of the notes by virtue of any such conflict.

For purposes of the change of control offer provisions of the notes, the following terms will be applicable:

“*Beneficial owner*” shall be determined in accordance with Rules 13d-3 and 13d-5 under the Exchange Act or any successor provisions, except that a person will be deemed to have beneficial ownership of all shares that person has the right to acquire irrespective of whether that right is exercisable immediately or only after the passage of time.

“*Change of control*” means the occurrence of any of the following: (1) the consummation of any transaction (including, without limitation, any merger or consolidation) the result of which is that any person or group (other than our company or one of our subsidiaries) becomes the beneficial owner, directly or indirectly, of more than 50% of our voting stock or other voting stock into which our voting stock is reclassified, consolidated, exchanged or changed, measured by voting power rather than number of shares; provided, however, that a person shall not be deemed beneficial owner of, or to own beneficially, (A) any securities tendered pursuant to a tender or exchange offer made by or on behalf of such person or any of such person’s affiliates until such tendered securities are accepted for purchase or exchange thereunder, or (B) any securities if such beneficial ownership (i) arises solely as a result of a revocable proxy delivered in response to a proxy or consent solicitation made pursuant to the applicable rules and regulations under the Exchange Act, and (ii) is not also then reportable on Schedule 13D (or any successor schedule) under the Exchange Act; (2) the direct or indirect sale, transfer, conveyance or other disposition (other than by way of merger or consolidation), in one or more series of related transactions, of all or substantially all of our assets and the assets of our subsidiaries, taken as a whole, to one or more persons or groups (other than our company or one of our subsidiaries), provided that none of the circumstances in this clause (2) will be a change of control if the persons that beneficially own our voting stock immediately prior to the transaction own, directly or indirectly, shares with a majority of the total voting power of all outstanding voting securities of the surviving or transferee person that are entitled to vote generally in the election of that person’s board of directors, managers or trustees immediately after the transaction; (3) we consolidate with, or merge with or into any person, or any person consolidates with, or merges with or into, us, in any such event pursuant to a transaction in which any of our outstanding voting stock or the voting stock of such other person is converted into or exchanged for cash, securities or other property, other than such transaction where the shares of our voting stock outstanding immediately prior to

such transaction constitute, or are converted into or exchanged for, a majority of the voting stock of the surviving person or any direct or indirect parent company of the surviving person immediately after giving effect to such transaction; or (4) the adoption of a plan relating to our liquidation or dissolution. Notwithstanding the foregoing, a transaction will not be deemed to involve a change of control under clause (1) above if (i) we become a direct or indirect wholly-owned subsidiary of a holding company and (ii) (A) the direct or indirect holders of the voting stock of such holding company immediately following that transaction are substantially the same as the holders of our voting stock immediately prior to that transaction or (B) immediately following that transaction no person (other than a holding company satisfying the requirements of this sentence) is the beneficial owner, directly or indirectly, of more than 50% of the voting stock of such holding company.

“*Change of control triggering event*” means the occurrence of both a change of control and a rating event.

“*Fitch*” means Fitch, Inc., and its successors.

“*Group*” has the meaning given by Section 13(d) and 14(d) of the Exchange Act or any successor provisions and includes any group acting for the purpose of acquiring, holding or disposing of securities within the meaning of Rule 13d-5(b)(1) under the Exchange Act or any successor provision.

“*Investment grade rating*” means a rating equal to or higher than Baa3 (or the equivalent) by Moody’s, BBB—(or the equivalent) by S&P and BBB—(or the equivalent) by Fitch, and the equivalent investment grade credit rating from any additional rating agency or rating agencies selected by us.

“*Moody’s*” means Moody’s Investors Service, Inc., and its successors.

“*Person*” has the meaning given by Section 13(d) and 14(d) of the Exchange Act or any successor provisions.

“*Rating agencies*” means (1) each of Fitch, Moody’s and S&P; and (2) if any of Fitch, Moody’s or S&P ceases to rate the notes or fails to make a rating of the notes publicly available for reasons outside of our control, a “nationally recognized statistical rating organization” within the meaning of Section 3(a)(62) of the Exchange Act selected by us (as certified by a resolution of our Board of Directors) as a replacement agency for Fitch, Moody’s or S&P, or all of them, as the case may be.

“*Rating event*” means the rating on the applicable series of notes is lowered by at least two of the three rating agencies and the notes are rated below an investment grade rating by at least two of the three rating agencies on any day during the period commencing 60 days prior to the first public notice of the occurrence of a change of control or our intention to effect a change of control and ending 60 days following consummation of such change of control (which period will be extended so long as the rating of the applicable series of notes is under publicly announced consideration for a possible downgrade by any of the rating agencies).

“*S&P*” means Standard & Poor’s Rating Services, a division of The McGraw-Hill Companies, Inc., and its successors.

“*Voting stock*” as applied to stock of any person, means shares, interests, participations or other equivalents in the equity interest (however designated) in such person having ordinary voting power for the election of a majority of the directors (or the equivalent) of such person, other than shares, interests, participations or other equivalents having such power only by reason of the occurrence of a contingency.

Certain Covenants

Limitation on Liens

We will not, nor will we permit any of our Subsidiaries to, create or incur any Lien on any of our or their respective Properties, whether now owned or hereafter acquired, or upon any income or profits therefrom, in order to secure any of our Indebtedness, without effectively providing that each series of notes shall be equally and ratably secured until such time as such Indebtedness is no longer secured by such Lien, except:

- (1) Liens existing as of the first issue date of the notes;
- (2) Liens granted after the first issue date of the notes on any of our or our Subsidiaries' Properties securing our Indebtedness created in favor of the holders of the notes;
- (3) Liens securing our Indebtedness which are incurred to extend, renew or refinance Indebtedness which is secured by Liens permitted to be incurred under the Indenture; provided that those Liens do not extend to or cover any of our or our Subsidiaries' Property other than the Property securing the Indebtedness being refinanced and that the principal amount of such Indebtedness does not exceed the principal amount of the Indebtedness being refinanced;
- (4) Liens created in substitution of or as replacements for any Liens permitted by the clauses directly above, provided that, based on a good faith determination of one of our officers, the Property encumbered under any such substitute or replacement Lien is substantially similar in nature to the Property encumbered by the otherwise permitted Lien which is being replaced; and
- (5) Permitted Liens.

Notwithstanding the foregoing, we and any of our Subsidiaries may, without securing any series of notes, create or incur Liens which would otherwise be subject to the restrictions set forth in the preceding paragraph, if after giving effect thereto, Exempted Debt does not exceed the greater of (a) 35% of Consolidated Net Worth calculated as of the date of the creation or incurrence of the Lien or (b) 35% of Consolidated Net Worth calculated as of the first issue date of the notes.

Limitation on Sale and Lease-Back Transactions

We will not, nor will we permit any of our Subsidiaries to, enter into any sale and lease-back transaction for the sale and leasing back of any Property, whether now owned or hereafter acquired, of ours or any of our Subsidiaries, unless:

- (1) such transaction was entered into prior to the first issue date of the notes;
- (2) such transaction was for the sale and leasing back to us of any Property by one of our Subsidiaries;
- (3) such transaction involves a lease for less than three years;
- (4) we would be entitled to incur Indebtedness secured by a mortgage on the property to be leased in an amount equal to the Attributable Liens with respect to such sale and lease-back transaction without equally and ratably securing the notes pursuant to the first paragraph of "—Limitation on Liens" above; or
- (5) we apply an amount equal to the fair value of the Property sold to the purchase of Property or to the retirement of our or any of our Subsidiaries' long-term Indebtedness within 120 days of the effective date of any such sale and lease-back transaction. In lieu of applying such amount to such retirement, we may, or may cause any of our Subsidiaries to, deliver debt securities to the Trustee therefor for cancellation, such debt securities to be credited at the cost thereof to us.

Notwithstanding the foregoing, we and any of our Subsidiaries may enter into any sale lease-back transaction which would otherwise be subject to the foregoing restrictions if after giving effect thereto and at the time of determination, Exempted Debt does not exceed the greater of (a) 35% of Consolidated Net Worth calculated as of the closing date of the sale-leaseback transaction or (b) 35% of Consolidated Net Worth calculated as of the first issue date of the notes.

Events of Default

Event of default means, with respect to each series of notes, any of the following:

- default in the payment of any interest on the notes of that series when it becomes due and payable, and continuance of such default for a period of 30 days (unless the entire amount of the payment is deposited by us with the Trustee or with the principal paying agent prior to the expiration of the 30-day period);
- default in the payment of principal of the notes of that series at their maturity;
- default in the performance or breach of any other covenant or warranty by us in the Indenture (other than defaults pursuant to the previous two bullet points above or pursuant to a covenant or warranty that has been included in the Indenture solely for the benefit of a series of debt securities other than that series of notes), which default continues uncured for a period of 90 days after we receive written notice from the Trustee or we and the Trustee receive written notice from the holders of not less than a majority in principal amount of the outstanding Notes of the affected series as provided in the Indenture; or
- certain voluntary or involuntary events of bankruptcy, insolvency or reorganization of our company.

No event of default with respect to the Notes (except as to certain events of bankruptcy, insolvency or reorganization) necessarily constitutes an event of default with respect to any other series of debt securities. The occurrence of an event of default may constitute an event of default under our bank credit agreements in existence from time to time. In addition, the occurrence of certain events of default or an acceleration under the Indenture may constitute an event of default under certain of our other indebtedness outstanding from time to time.

We will provide the Trustee written notice of any default or event of default within 30 days of becoming aware of the occurrence of such default or event of default, which notice will describe in reasonable detail the status of such default or event of default and what action we are taking or propose to take in respect thereof.

If an event of default with respect to a series of Notes occurs and is continuing (other than an event of default regarding certain events of bankruptcy, insolvency or reorganization of our company), then the Trustee or the holders of not less than a majority in principal amount of the outstanding Notes of that series may, by a notice in writing to us (and to the Trustee if given by the holders), declare to be due and payable immediately the principal of, and accrued and unpaid interest, if any, on all Notes of that series. In the case of an event of default resulting from certain events of bankruptcy, insolvency or reorganization, the principal of and accrued and unpaid interest, if any, on all outstanding debt securities issued under the Indenture will become and be immediately due and payable without any declaration or other act on the part of the Trustee or any holder of outstanding debt securities, including the Notes. At any time after a declaration of acceleration with respect to a series of Notes has been made, and before a judgment or decree for payment of the money due has been obtained by the Trustee, the holders of a majority in principal amount of the outstanding Notes of that series may, by written notice to us and the Trustee, rescind and annul such acceleration if all events of default, other than the non-payment of accelerated principal and interest, if any, with respect to the Notes of that series, have been cured or waived as provided in the Indenture.

The Indenture provides that the Trustee will be under no obligation to exercise any of its rights or powers under the Indenture at the request of any holder of notes, unless the Trustee receives indemnity satisfactory to it against any cost, liability or expense which might be incurred by it in exercising such right or power. Subject to certain rights of the Trustee, the holders of a majority in principal amount of the outstanding Notes of the affected series will have the right to direct the time, method and place of conducting any proceeding for any remedy available to the Trustee or exercising any trust or power conferred on the Trustee with respect to the Notes of that series.

No holder of any Note of any series will have any right to institute any proceeding, judicial or otherwise, with respect to the Indenture, or for the appointment of a receiver or Trustee, or for any remedy under the Indenture unless, among other things:

- that holder has previously given to the Trustee written notice of a continuing event of default with respect to the Notes of that series; and

- the holders of at least a majority in principal amount of the outstanding Notes of that series have made written request, and offered reasonable indemnity or security, to the Trustee to institute the proceeding as Trustee, and the Trustee has not received from the holders of a majority in principal amount of the outstanding Notes of that series a direction inconsistent with that request and has failed to institute the proceeding within 60 days.

Notwithstanding any other provision in the Indenture, the holder of any Note will have an absolute and unconditional right to receive payment of the principal of, premium and any interest on that Note on or after the due dates expressed in that Note and to institute suit for the enforcement of any such payment.

If any securities are outstanding under the Indenture, the Indenture requires us, within 120 days after the end of each fiscal year, to furnish to the Trustee a statement as to our compliance with the indenture. If a default or event of default occurs and is continuing with respect to notes of any series and if it is known to a responsible officer of the Trustee, the Trustee shall deliver to each holder of the Notes of that series notice of a default or event of default within 90 days after it occurs. The Indenture provides that the Trustee may withhold notice to the holders of the Notes of any default or event of default (except in the case of a default or event of default in payment of principal of or interest on any Note of that series) with respect to Notes of that series if it in good faith determines that withholding notice is in the interest of the holders of those Notes.

Modification and Waiver

We and the Trustee may modify and amend the Indenture or Notes of any series without the consent of any holder of Notes:

- to cure any ambiguity, defect or inconsistency;
- to comply with the covenant described below under the heading “—Consolidation, Merger and Sale of Assets;”
- to provide for uncertificated notes in addition to or in place of certificated notes;
- to add guarantees with respect to Notes of any series or secure notes of any series;
- to surrender any of our rights or powers under the Indenture;
- to add covenants or events of default for the benefit of the holders of Notes of any series;
- to comply with the applicable procedures of the applicable depositary;
- to make any change that would not adversely affect the rights of any holder of Notes in any material respect;
- to provide for the issuance of and establish the form and terms and conditions of additional Notes of any series as permitted by the Indenture;
- to effect the appointment of a successor trustee with respect to the Notes and to add to or change any of the provisions of the Indenture to provide for or facilitate administration by more than one trustee; or
- to comply with requirements of the U.S. Securities and Exchange Commission in order to effect or maintain the qualification of the Indenture under the U.S. Trust Indenture Act of 1939.

We may also modify and amend the Indenture with the consent of the holders of at least a majority in principal amount of the outstanding Notes of each series affected by the modifications or amendments. We may not

make any modification or amendment without the consent of the holders of each affected Note then outstanding if that amendment will:

- reduce the amount of Notes whose holders must consent to an amendment, supplement or waiver;
- reduce the rate of or extend the time for payment of interest (including any additional amounts) on the Notes;
- reduce the principal of or premium on or change the fixed maturity of the Notes;
- waive a default in the payment of the principal of, premium or interest on the notes (except a rescission of acceleration of the notes by the holders of at least a majority in aggregate principal amount of the then outstanding Notes of that series and a waiver of the payment default that resulted from such acceleration);
- make the principal of or interest on the Notes payable in currency other than that stated in the Notes;
- make any change to certain provisions of the Indenture relating to, among other things, the right of holders of the Notes to receive payment of the principal of, premium and interest on the Notes and to institute suit for the enforcement of any such payment and to waivers or amendments; or
- waive a redemption payment with respect to the Notes.

Except for certain specified provisions, the holders of at least a majority in principal amount of the outstanding Notes of the affected series may, on behalf of the holders of all the Notes of that series, waive our compliance with provisions of the Indenture. The holders of a majority in principal amount of the outstanding Notes of the affected series may, on behalf of the holders of all the Notes of such series, waive any past default under the Indenture with respect to that series and its consequences, except a default in the payment of the principal of, premium or any interest on any Note of that series; provided, however, that the holders of a majority in principal amount of the outstanding Notes of the affected series may rescind an acceleration and its consequences, including any related payment default that resulted from such acceleration.

No amendment to cure any ambiguity, defect or inconsistency in the Indenture made solely to conform the Indenture to the description of notes contained in the applicable prospectus supplement will be deemed to adversely affect the interests of the holders of the Notes.

Consolidation, Merger and Sale of Assets

We may not consolidate with or merge with or into, or convey, transfer or lease all or substantially all of our properties and assets to, any person, which we refer to as a “successor person,” unless:

- we are the surviving corporation or the successor person (if other than Amgen) is organized and validly existing under the laws of any U.S. domestic jurisdiction and expressly assumes, pursuant to a supplemental Indenture, our obligations on the notes and under the Indenture; and
- immediately after giving effect to the transaction, no default or event of default shall have occurred and be continuing under the Indenture.

Notwithstanding the foregoing, any of our Subsidiaries may consolidate with, merge into or transfer all or part of its properties and assets to us.

Defeasance and Covenant Defeasance

Legal Defeasance

The Indenture provides that we may be discharged from any and all obligations in respect of the Notes (subject to certain exceptions). We will be so discharged upon the deposit with the Trustee, in trust, of money, U.S. government obligations and/or foreign government obligations that, through the payment of interest and principal in accordance with their terms, will provide money, U.S. government obligations or foreign government obligations in an amount sufficient in the opinion of a nationally recognized firm of independent public accountants or investment bank to pay and discharge each installment of principal of, premium and interest on the Notes on the stated maturity of those payments in accordance with the terms of the Indenture and the Notes.

This discharge may occur only if, among other things, we have delivered to the Trustee an opinion of counsel stating that we have received from, or there has been published by, the IRS a ruling or, since the date of execution of the Indenture, there has been a change in the applicable U.S. federal income tax law, in either case to the effect that, and based thereon such opinion shall confirm that, the holders of the Notes will not recognize income, gain or loss for U.S. federal income tax purposes as a result of the deposit, defeasance and discharge and will be subject to U.S. federal income tax on the same amounts and in the same manner and at the same times as would have been the case if such deposit, defeasance and discharge had not occurred.

Defeasance of Certain Covenants

The Indenture provides that upon compliance with certain conditions:

- we may omit to comply with the covenant described under the heading “—Consolidation, Merger and Sale of Assets” and certain other covenants set forth in the Indenture, as well as any additional covenants set forth in the applicable prospectus supplement; and
- any omission to comply with those covenants will not constitute a default or an event of default with respect to the Notes, which we refer to as a “covenant defeasance.”

The conditions include:

- depositing with the Trustee money, U.S. government obligations and/or foreign government obligations that, through the payment of interest and principal in accordance with their terms, will provide money in an amount sufficient in the opinion of a nationally recognized firm of independent public accountants or investment bank to pay and discharge each installment of principal of, premium and interest on the notes on the stated maturity of those payments in accordance with the terms of the Indenture and the Notes; and
- delivering to the Trustee an opinion of counsel to the effect that the holders of the Notes will not recognize income, gain or loss for U.S. federal income tax purposes as a result of the deposit and related covenant defeasance and will be subject to U.S. federal income tax on the same amounts and in the same manner and at the same times as would have been the case if the deposit and related covenant defeasance had not occurred.

Covenant Defeasance and Events of Default

In the event we exercise our option to effect covenant defeasance with respect to any series of the Notes and the Notes of that series are declared due and payable because of the occurrence of any event of default, the amount of money, U.S. government obligations and/or foreign government obligations on deposit with the Trustee will be sufficient to pay amounts due on the Notes of that series at the time of their stated maturity but may not be sufficient to pay amounts due on the notes of that series at the time of the acceleration resulting from the event of default. In such a case, we would remain liable for those payments.

Concerning the Trustee

The Bank of New York Mellon Trust Company, N.A. is Trustee under the Indenture.

Governing Law

The Indenture and the Notes, including any claim or controversy arising out of or relating to the Indenture or the Notes, are governed by the laws of the State of New York.

Form of Award Notice

[The information set forth in this Award Notice will be contained on the related pages on Merrill Lynch Benefits Website (or the website of any successor company to Merrill Lynch Bank & Trust Co., FSB). This Award Notice shall be replaced by the equivalent pages on such website. References to Award Notice in this Agreement shall then refer to the equivalent pages on such website.]

This notice of Award (the “Award Notice”) sets forth certain details relating to the grant by the Company to you of the Award identified below, pursuant to the Plan. The terms of this Award Notice are incorporated into the Stock Option Agreement (the “Agreement”) that accompanies this Award Notice and made part of the Agreement. Capitalized terms used in this Award Notice that are not otherwise defined in this Award Notice have the meanings given to such terms in the Agreement.

Employee:

Employee ID:

Address:

Award Type:

Grant ID:

Plan: Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan, as amended and/or restated from time to time

Grant Date:

Grant Price: \$ _____

Number of Shares

Covered by Option:

Type: Non-qualified Stock Option

Expiration Date: The [_____] (____th) anniversary of the Grant Date

Vesting Date: Means the vesting date indicated in the Vesting Schedule

Vesting Schedule: Means the schedule of vesting set forth under Vesting Details

Vesting Details: Means the presentation (tabular or otherwise) of the Vesting Date and the quantity of Shares vesting.

IMPORTANT NOTICE REGARDING ACCEPTANCE OF THE AWARD AND THE REQUIREMENT TO OPEN A BROKERAGE ACCOUNT:

RESIDENTS OF THE U.S. AND PUERTO RICO: Please read this Award Notice, the Plan and the Agreement (collectively, the “Grant Documents”) carefully. If you, as a resident of the U.S. or Puerto Rico, do **not** wish to receive this Award and/or you do **not** consent and agree to the terms and conditions on which this Award is offered, as set forth in the Grant Documents, then you must reject the Award by contacting the Merrill Lynch call center at +1 (800) 97AMGEN (+1 (800) 972-6436) within the U.S., Puerto Rico and Canada or +1 (609) 818-8910

ⁱ This provision is only for use on the form of grant used for the U.S. and Puerto Rico.

from all other countries (Merrill Lynch will accept the charges for your call) no later than the forty-fifth calendar day following the day on which this Award Notice is made available to you, in which case the Award will be cancelled. For the purpose of determining the forty-five calendar days, Day 1 will be the day **immediately** following the day on which this Award Notice is made available to you. Your failure to notify the Company of your rejection of the Award or your refusal of, or disagreement with, all terms and conditions of the Award, as set forth in the Grant Documents, within this specified period will constitute your acceptance of the Award and your agreement with all terms and conditions of the Award, as set forth in the Grant Documents. If you agree to the terms and conditions of your grant and you desire to accept it, then no further action is needed on your part to accept the grant. However, you must still open a brokerage account as directed by the Company, by 1:00 pm Pacific Time on or before the date that is 11 months after the Grant Date specified in this Award Notice. This step is necessary to process transactions related to your equity grant. **If you do not open a brokerage account by this deadline, your grant will be cancelled.**

GRANT OF STOCK OPTION AGREEMENT

THE SPECIFIC TERMS OF YOUR STOCK OPTION ARE FOUND IN THE PAGES RELATING TO THE GRANT OF STOCK OPTIONS FOUND ON MERRILL LYNCH BENEFITS WEBSITE (OR THE WEBSITE OF ANY SUCCESSOR COMPANY TO MERRILL LYNCH BANK & TRUST CO., FSB) (THE “AWARD NOTICE”) WHICH ACCOMPANIES THIS DOCUMENT. THE TERMS OF THE AWARD NOTICE ARE INCORPORATED INTO THIS GRANT OF STOCK OPTIONS.

On the Grant Date, specified in the Award Notice, Amgen Inc., a Delaware corporation (the “Company”), has granted to you, the grantee named in the Award Notice, under the plan specified in the Award Notice (the “Plan”), an option (the “Option”) to purchase the number of shares of the \$0.0001 par value common stock of the Company (the “Shares”) specified in the Award Notice, pursuant to the terms set forth in this Stock Option Agreement, any additional terms and conditions for your country set forth in the attached Appendix A and the Award Notice (collectively, the “Agreement”) and the Plan. This Option is not intended to qualify and will not be treated as an “incentive stock option” within the meaning of Section 422 of the U.S. Internal Revenue Code of 1986, as amended (together with the regulations and other official guidance promulgated thereunder, the “Code”). Capitalized terms not defined herein shall have the meanings assigned to such terms in the Plan.

The terms and conditions of your Option are as follows:

I. Subject to the terms and conditions of the Plan and this Agreement, on each Vesting Date the Option shall vest with respect to the number of Shares indicated on the Vesting Schedule, provided that you have remained continuously and actively employed with the Company or an Affiliate through each applicable Vesting Date, unless [(i) your employment has terminated due to your Voluntary Termination (as defined in Section IV(A)(5)) or (ii)]*ⁱⁱ you experience a Qualified Termination (as defined in Section IV(B)(4)), or as otherwise determined by the Company in the exercise of its discretion as provided in Section IV(A)(7). This Option may only be exercised for whole shares of the Common Stock, and the Company shall be under no obligation to issue any fractional Shares to you. Subject to the limitations contained herein, this Option shall be exercisable with respect to each installment on or after the applicable Vesting Date. Notwithstanding anything herein to the contrary, the Vesting Schedule may be accelerated (by notice in writing) by the Company in its sole discretion at any time during the term of this Option. In addition, if not prohibited by local law, vesting may be suspended by the Company in its sole discretion during a leave of absence as provided from time to time according to Company policies and practices; provided, that, in no event shall any such suspension extend the term of this Option beyond the Expiration Date set forth on the Award Notice and in this Agreement.

ⁱⁱ *Section IV(A)(5) of this Agreement is not applicable to awards identified by the Administrator as new hire, retention or promotion grants and the provisions of such section shall be reserved and references thereto identified by an asterisk (*) shall be omitted from the agreements evidencing such grants.*

II. (1) The per Share exercise price of this Option is the Grant Price as defined in the Award Notice, being not less than the Fair Market Value of the Common Stock on the Grant Date of this Option.

(2) To the extent permitted by applicable statutes and regulations, payment of the exercise price per Share is due in full upon exercise of all or any part of each installment which has become exercisable by you by means of (i) cash or a check, (ii) any cashless exercise procedure through the use of a brokerage arrangement approved by the Company, or (iii) any other form of legal consideration that may be acceptable to the Board or the Committee in their discretion.

(3) Notwithstanding anything in Section II(2), to the extent permitted by applicable statutes and regulations, if, at the time of exercise, the Company's Common Stock is publicly traded and quoted regularly in the Wall Street Journal, payment of the exercise price may be made by delivery of already-owned Shares with a Fair Market Value equal to the exercise price of the Shares for which this Option is being exercised. The already-owned Shares must have been owned by you for the period required to avoid adverse accounting treatment and owned free and clear of any liens, claims, encumbrances or security interests. Payment may also be made by a combination of cash and already-owned Shares.

Notwithstanding the foregoing, the Company reserves the right to restrict the methods of payment of the exercise price if necessary or advisable to comply with applicable law or regulation, as determined by the Company in its sole discretion.

III. Notwithstanding anything to the contrary contain herein, the Company shall not take any actions that would violate the Securities Act, the Exchange Act, the Code, or any other securities or tax or other applicable law or regulation, or the rules of any Securities Exchange. The Company, in its sole discretion, may impose any timing or other restrictions with respect to the exercise of this Option arising from compliance with any securities or tax laws or other rules or regulations. Notwithstanding anything to the contrary contained herein, this Option may not be exercised and no Shares underlying the Option will be issued unless such Shares are then registered under the Securities Act, or, if such Shares are not then so registered, the Company has determined that such exercise and issuance would be exempt from the registration requirements of the Securities Act, and that the issuance satisfies all other applicable legal requirements. If the Option cannot be exercised and expires during this period, you will forfeit the Option and no Shares or value will be transferred to you.

IV. (A) The term of this Option commences on the Grant Date and, unless sooner terminated as set forth below or in the Plan, terminates on the [_____] (___th) anniversary of the Grant Date (the "Expiration Date"). This Option shall terminate prior to the Expiration Date as follows: three (3) months after the termination of your employment with the Company or an Affiliate (as defined in the Plan) for any reason or for no reason, including if your employment is terminated by the Company or an Affiliate without Cause (as defined below), or in the event of

any other termination of your employment caused directly or indirectly by the Company or an Affiliate, unless:

(1) such termination of your employment is due to your Permanent and Total Disability (as defined below), in which case (i) the Option shall terminate on the Expiration Date and (ii) the vesting of the Option shall be accelerated in full and the Option shall be fully exercisable, subject to your execution and non-revocation of a general release and waiver in a form provided by the Company (for the purpose of resolving any potential or actual disputes arising from your employment and the termination of your employment with the Company) (a “Release”) as of the day immediately preceding such termination of your employment with respect to the Option. Notwithstanding the foregoing, if the Option was granted in the calendar year in which such termination occurs, (i) the Option shall instead be accelerated to vest only with respect to a number of Shares equal to (A) the number of Shares subject to the Option multiplied by (B) a fraction, the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12), subject to your execution and non-revocation of a Release, and (ii) any portion of the Option (if any) that remains unvested following the acceleration provided for in clause (i) shall automatically expire and terminate on the date of the termination of your active employment due to your Permanent and Total Disability without consideration therefor;

(2) such termination of your employment is due to your death, in which case (i) the Option shall terminate on the Expiration Date and (ii) the vesting of the Option shall be accelerated to vest in full and the Option shall be fully exercisable as of the day immediately preceding your death with respect to the Option. Notwithstanding the foregoing, if the Option was granted in the calendar year in which your death occurs (i) the Option shall instead be accelerated to vest only with respect to a number of Shares equal to (A) the number of Shares subject to the Option multiplied by (B) a fraction, the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12), and (ii) any portion of the Option (if any) that remains unvested following the acceleration provided for in clause (i) shall automatically expire and terminate on the date of termination of your active employment due to your death without consideration therefor;

(3) during any part of such three (3) month period, this Option is not exercisable solely because of the condition set forth in Section III above, in which event this Option shall not terminate until the earlier of the Expiration Date or until it shall have been exercisable for an aggregate period of three (3) months after the termination of your employment;

(4) exercise of this Option within three (3) months after termination of your employment with the Company or with an Affiliate would result in liability under Section 16(b) of the Exchange Act, in which case this Option will terminate on the earliest of: (a) the tenth (10th) day after the last date upon which exercise would result in such liability; (b) six (6) months and ten (10) days after the termination of your employment with the Company or an Affiliate; or (c) the Expiration Date;

(5) [such termination of your employment is due to your voluntary termination (and such voluntary termination is not the result of Permanent and Total Disability (as defined below)) after you are at least sixty five (65) years of age, or after you are at least fifty-five (55) years of age and have been an employee of the Company and/or an Affiliate for at least ten (10) years in the aggregate as determined by the Company in its sole discretion according to Company policies and practices as in effect from time to time (“Voluntary Termination”), in which case (i) this Option shall terminate on the Expiration Date and (ii) the unvested portions of this Option will become exercisable pursuant to the Vesting Schedule without regard to your Voluntary Termination of your employment, subject to your execution and non-revocation of a Release. Notwithstanding the foregoing, if the Option was granted in the calendar year in which your Voluntary Termination occurs, (i) the Option will continue to vest and become exercisable pursuant to the Vesting Schedule only with respect to (A) a number of Shares equal to the number of Shares subject to the Option multiplied by (B) a fraction, the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12), and (ii) any portion of the Option (if any) that remains unvested following the acceleration provided for in clause (i) shall automatically expire and terminate on the date of the termination of your active employment due to your Voluntary Termination without consideration therefor. Notwithstanding the foregoing, in the event your Voluntary Termination occurs on or after the date of a Change of Control, then, to the extent permitted by applicable law, the vesting of the Option shall be accelerated to vest in full as of the day immediately prior to the date of your Voluntary Termination. Notwithstanding the definition of Voluntary Termination set forth above, if the Company receives an opinion of counsel that there has been a legal judgment and/or legal development in your jurisdiction that would likely result in the favorable treatment upon Voluntary Termination described above being deemed unlawful and/or discriminatory, then the Committee will not apply the favorable treatment described above;][Reserved]*ⁱⁱⁱ

(6) such termination of your employment is due to a Qualified Termination, in which case, the Option shall terminate on the earlier of (a) the date that is three (3) months following the date of such Qualified Termination or (b) the Expiration Date, and, to the extent permitted by applicable law, the vesting of the Option shall be accelerated and the Option shall be fully exercisable as of the day immediately prior to the Qualified Termination; or

(7) the Company determines, in its sole discretion at any time during the term of this Option, in writing, to otherwise extend the period of time during which this Option will vest and may be exercised after termination of your employment; provided, that, in no event shall any such extension extend the term of this Option beyond the Expiration Date set forth on the Award Notice and in this Agreement.

ⁱⁱⁱ Section IV(A)(5) of this Agreement is not applicable to awards identified by the Administrator as new hire, retention or promotion grants and the provisions of such section shall be reserved and references thereto identified by an asterisk (*) shall be omitted from the agreements evidencing such grants.

However, in any and all circumstances and except to the extent the Vesting Schedule has been accelerated by the Company in its sole discretion during the term of this Option or as a result of your Permanent and Total Disability or death as provided in Sections IV(A)(1) or IV(A)(2) above, respectively, [as a result of your Voluntary Termination as provided in Section IV(A)(5) above,]* as a result of a Qualified Termination as provided in Section IV(A)(6) above or as otherwise determined by the Company in the exercise of its discretion as provided in Section IV(A)(7) above, this Option may be exercised following termination of your employment only as to that number of Shares as to which it was exercisable on the date of termination of your employment under the provisions of Section I of this Agreement.

(B) For purposes of this Option:

(1) “termination of your employment” shall mean the last date you are either an active employee of the Company or an Affiliate or actively engaged as a Director to the Company or an Affiliate; in the event of termination of your employment (whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any), your right to receive options and vest under the Plan, if any, will terminate effective as of the date that you are no longer actively employed and will not be extended by any notice period (e.g., active employment would not include any period of “garden leave” or similar period mandated under employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any). Your right, if any, to exercise the Option after termination of employment will be measured by the date of termination of your active employment and will not be extended by any notice period mandated under local law. The Administrator shall have the exclusive discretion to determine when you are no longer actively providing services for purposes of this Agreement (including whether you may still be considered to be providing services while on a leave of absence);

(2) “Cause” shall mean (i) your conviction of a felony (or similar crime under applicable law, as determined by the Company), or (ii) your engaging in conduct that constitutes willful gross neglect or willful gross misconduct in carrying out your duties, resulting, in either case, in material economic harm to the Company or any Affiliate, unless you believed in good faith that such conduct was in, or not contrary to, the best interests of the Company or any Affiliate. For purposes of clause (ii) above, no act, or failure to act, on your part shall be deemed “willful” unless done, or omitted to be done, by you not in good faith;

(3) “Permanent and Total Disability” shall have the meaning ascribed to such term under Section 22(e)(3) of the Code and with such permanent and total disability being certified prior to termination of your employment by (a) the U.S. Social Security Administration, (b) the comparable governmental authority applicable to an Affiliate, (c) such other body having the relevant decision-making power applicable to an Affiliate, or (d) an independent medical advisor appointed by the Company in its sole discretion, as applicable, in any such case;

(4) “Qualified Termination” shall mean

(a) if you are an employee who participates in the Change of Control Plan (as defined below), your termination of employment within two (2) years following a Change of Control (i) by the

Company or an Affiliate other than for Cause, Disability (as defined below) or as a result of your death, or (ii) by you for Good Reason (as defined in the Change of Control Plan); or

(b) if you are an employee who does not participate in the Change of Control Plan or the Change of Control Plan is no longer in effect, your termination of employment within two (2) years following a Change of Control by the Company or an Affiliate other than for Cause, Disability (as defined below) or as a result of your death;

(5) “Change of Control” shall mean the occurrence of any of the following:

(a) the acquisition (other than from the Company) by any person, entity or “group,” within the meaning of Section 13(d)(3) or 14(d)(2) of the Exchange Act (excluding, for this purpose, the Company or any of its Affiliates, or any employee benefit plan of the Company or any of its Affiliates which acquires beneficial ownership of voting securities of the Company), of beneficial ownership (within the meaning of Rule 13d-3 promulgated under the Exchange Act) of fifty percent (50%) or more of either the then outstanding Shares or the combined voting power of the Company’s then outstanding voting securities entitled to vote generally in the election of directors; or

(b) the consummation by the Company of a reorganization, merger, consolidation, (in each case, with respect to which persons who were the stockholders of the Company immediately prior to such reorganization, merger or consolidation do not, immediately thereafter, own more than fifty percent (50%) of the combined voting power entitled to vote generally in the election of directors of the reorganized, merged or consolidated company’s then outstanding voting securities) or a liquidation or dissolution of the Company or of the sale of all or substantially all of the assets of the Company.

Notwithstanding anything herein or in any Award Agreement to the contrary, if a Change of Control constitutes a payment event with respect to any Award that is subject to United States income tax and which provides for a deferral of compensation that is subject to Section 409A of the Code, the transaction or event described in subsection (a) or (b), (c) above must also constitute a “change in control event,” as defined in U.S. Treasury Regulation §1.409A-3(i)(5), in order to constitute a Change of Control for purposes of payment of such Award.

(6) “Change of Control Plan” shall mean the Company’s change of control and severance plan, including the Amgen Inc. Change of Control Severance Plan, as amended and restated, effective as of December 9, 2010 (and any subsequent amendments thereto), or any equivalent plan governing the provision of benefits to eligible employees upon the occurrence of a Change of Control (including resulting from a termination of employment that occurs within a specified time period following a Change of Control), as in effect immediately prior to a Change of Control; and

(7) “Disability” shall mean your “disability” as determined in accordance with the Company’s long-term disability plan as in effect immediately prior to a Change of Control.

V. (A) To the extent specified above, this Option may be exercised by delivering a notice of exercise in person, by mail, via electronic mail or facsimile or by other authorized method designated by the Company, together with the exercise price to the Company Stock Administrator, or to such other person as the Company Stock Administrator may designate,

during regular business hours, together with such additional documents as the Company may then require pursuant to Section 7.2(b) of the Plan.

(B) Regardless of any action the Company or your actual employer (the “Employer”) takes with respect to any or all income tax (including federal, state and local taxes), social insurance, payroll tax, fringe benefit tax, payment on account or other tax-related items related to your participation in the Plan and legally applicable to you (“Tax Obligations”), you acknowledge that the ultimate liability for all Tax Obligations is and remains your responsibility and may exceed the amount, if any, actually withheld by the Company and/or your Employer. You further acknowledge that the Company and/or your Employer: (a) make no representations or undertakings regarding the treatment of any Tax Obligations in connection with any aspect of the Option grant or the underlying Shares, including, but not limited to, the grant, vesting or exercise of the Option, the subsequent sale of Shares acquired pursuant to such exercise and the receipt of any dividends; and (b) do not commit to and are under no obligation to structure the terms of the grant or any aspect of the Option to reduce or eliminate your liability for Tax Obligations or achieve any particular tax result. Furthermore, if you become subject to tax in more than one jurisdiction, you acknowledge that the Company and/or your Employer (or former employer, as applicable) may be required to withhold or account for Tax Obligations in more than one jurisdiction.

(C) Prior to any relevant taxable or tax withholding event, as applicable, you shall pay or make adequate arrangements satisfactory to the Company and/or your Employer to satisfy all Tax Obligations. In this regard, you authorize the Company and/or your Employer, or their respective agents, at their discretion, to satisfy all applicable Tax Obligations by one or a combination of the following:

- (1) withholding from your wages or other cash compensation paid to you by the Company and/or your Employer;
- (2) withholding from proceeds of the sale of Shares acquired upon exercise of the Option either through your voluntary sale or through a mandatory sale arranged by the Company (on your behalf pursuant to this authorization); or
- (3) withholding in Shares issuable, or cash payable, upon exercise of the Option, provided that, if such Shares are withheld, the Company and your Employer shall only withhold an amount of Shares with a fair market value not to exceed the Tax Obligations as determined in the discretion of the Company or your Employer, as applicable.

Depending on the withholding method, the Company may withhold or account for Tax Obligations by considering applicable minimum statutory withholding rates or other applicable withholding rates, including maximum applicable rates. If the Tax Obligations are satisfied by withholding in Shares, for tax purposes you are deemed to have been issued the full number of Shares subject to the exercised Option, notwithstanding that a number of the Shares is

held back and not actually issued to you solely for the purpose of paying the Tax Obligations due as a result of any aspect of your participation in the Plan.

(D) Finally, you shall pay to the Company or your Employer any amount of Tax Obligations that the Company or your Employer may be required to withhold or account for as a result of your participation in the Plan that cannot be or were not satisfied by the means previously described. You agree to take any further actions and execute any additional documents as may be necessary to effectuate the provisions of this Section V. Notwithstanding anything to the contrary contained herein, the Company may refuse to issue or deliver the Shares or the proceeds of the sale of Shares if you fail to comply with your obligations in connection with the Tax Obligations.

VI. This Option is not transferable, except by will or the laws of descent and distribution, and is exercisable during your life only by you except if you have named a trust created for the benefit of you, your spouse, or members of your immediate family (a "Trust") as beneficiary of this Option, this Option may be exercised by the Trust after your death.

VII. Any notices provided for in this Option or the Plan shall be given in writing or electronically and shall be deemed effectively given upon receipt or, in the case of notices delivered by the Company to you, five (5) days after deposit in the United States mail or equivalent foreign postal service, postage prepaid, addressed to you at such address as is currently maintained in the Company's records or at such other address as you hereafter designate by written notice to the Company Stock Administrator. Such notices may be given using any automated system for the documentation, granting or exercise of Awards, such as a system using an internet website or interactive voice response, as approved by the Company.

VIII. This Option is subject to all the provisions of the Plan and its provisions are hereby made a part of this Option, including without limitation the provisions of Articles 6 and 7 of the Plan relating to Options, and is further subject to all interpretations, amendments, rules and regulations which may from time to time be promulgated and adopted pursuant to the Plan. In the event of any conflict between the provisions of this Option and those of the Plan, the provisions of the Plan shall control.

IX. In order for the Company to facilitate your participation in the Plan, the Company and your Employer must collect and use personal data about you. In accordance with applicable laws, reasonable security measures will be implemented and maintained to protect the security of your personal data; however, you understand that absolute security cannot be guaranteed.

You understand that the Company and your Employer may hold certain personal information about you, including your name, home address and telephone number, email address, date of birth, social insurance/security number (to the extent permitted under applicable local law), passport or other identification number, salary, nationality, job title/work history/service periods, residency status, citizenship, tax withholding and payroll data, any shares of stock or directorships held in the Company, details of all equity compensation or any other entitlement to Shares

awarded, cancelled, vested, unvested or outstanding in your favor, for the purposes of implementing, administering and managing the Plan (“personal data”).

You authorize the transfer of your personal data to Merrill Lynch Bank & Trust Co., FSB, or any successor thereto, and any other third parties which may assist the Company (presently or in the future) with implementing, administering and managing your participation in the Plan to receive, possess, use, retain and transfer your personal data, in electronic or other form, for the purpose of implementing, administering and managing your participation in the Plan, including any requisite transfer of such personal data as may be required to any other broker, escrow agent or other third party with whom the Shares received upon exercise of this Option may be deposited. You understand that such authorized recipients of your personal data may be located in countries that do not provide the same level of data privacy laws and protections as the country in which your personal data originated. Transfers of personal data among Company and its group entities follow applicable laws and our Binding Corporate Rules (BCRs). For more information on Company’s BCRs, please visit <http://www.amgen.com/bcr/>. You acknowledge that the collection, use and transfer of your personal data is necessary to facilitate your participation in the Plan, as well as to grant you Options or other equity awards and administer or maintain such awards.

You may correct or update your personal data previously provided to Company, by contacting your local human resources representative. Subject to applicable law, you may have additional rights, including the right to object and/or request destruction of your personal data. To exercise these rights, where applicable, please contact your local human resources representative.

X. The terms of this Option shall be governed by the laws of the State of Delaware without giving effect to principles of conflicts of laws. For purposes of litigating any dispute that arises hereunder, the parties hereby submit to and consent to the jurisdiction of the State of Delaware, and agree that such litigation shall be conducted in the courts of the State of Delaware, or the federal courts for the United States for the federal district located in the State of Delaware, and no other courts, where this Option is made and/or to be performed.

XI. Notwithstanding any provision of this Option to the contrary, if you are employed by the Company or an Affiliate in any of the countries identified in the attached Appendix A (which constitutes a part of this Agreement), are subject to the laws of any foreign jurisdiction, or relocate to one of the countries included in the attached Appendix A, the Option granted hereunder shall be subject to any additional terms and conditions for your country set forth in Appendix A and the following additional terms and conditions:

- a. the terms and conditions of this Option, including Appendix A, are deemed modified to the extent necessary or advisable to comply with applicable foreign laws or facilitate the administration to the Plan;
- b. if applicable, the effectiveness of this Option is conditioned upon its compliance with any applicable foreign laws, regulations, rules or

local governmental regulatory exemption and subject to receipt of any required foreign regulatory approvals; and

- c. the Company may take any other action before or after the date of this Option that it deems advisable to obtain approval or comply with any necessary local governmental regulatory exemptions or approvals.

XII. (A) In accepting this Option, you acknowledge, understand and agree that:

- (1) the Plan is established voluntarily by the Company, is discretionary in nature and may be modified, amended, suspended or terminated by the Company at any time, as provided in the Plan;
- (2) the Plan is operated and the Options are granted solely by the Company, and only the Company is a party to this Agreement; accordingly, any rights you may have under this Agreement may be raised only against the Company but not any Affiliate (including, but not limited to, the Employer);
- (3) no Affiliate (including, but not limited to, the Employer) has any obligation to make any payment of any kind to you under this Agreement;
- (4) the grant of this Option is exceptional, voluntary and occasional and does not create any contractual or other right to receive future awards of options, or benefits in lieu of options even if options have been awarded in the past;
- (5) all decisions with respect to future awards, if any, will be at the sole discretion of the Company;
- (6) your participation in the Plan is voluntary;
- (7) the grant of Options, the underlying Shares, and the income from and value of same, are not intended to replace any pension rights or compensation;
- (8) neither the grant of options nor any provision of this Option, the Plan or the policies adopted pursuant to the Plan confer upon you any right with respect to employment or continuation of current employment and shall not interfere with the ability of your Employer to terminate your employment or service relationship (if any) at any time;
- (9) in the event that you are not an employee of the Company or any Affiliate, the Option shall not be interpreted to form an employment contract or relationship with the Company or any Affiliate;
- (10) the future value of the underlying Shares is unknown, indeterminable, and cannot be predicted with certainty;

(11) if the underlying Shares do not increase in value, this Option will have no value; if you exercise this Option and obtain Shares, the value of those Shares acquired upon exercise may increase or decrease in value, even below the Grant Price per Share;

(12) in consideration of the grant of this Option, no claim or entitlement to compensation or damages arises from forfeiture of options resulting from (i) the termination of your employment by the Company or an Affiliate (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any) and/or (ii) the application of the Clawback Policy as described in Section XVIII below, and you irrevocably release the Company and your Employer from any such claim that may arise; if, notwithstanding the foregoing, any such claim is found by a court of competent jurisdiction to have arisen, you shall be deemed irrevocably to have waived your entitlement to pursue such claim;

(13) unless otherwise agreed with the Company, the Options, the underlying Shares, and the income from and value of same, are not granted as consideration for, or in connection with, the service you may provide as a director of an Affiliate of the Company;

(14) except as otherwise provided in this Agreement or the Plan, the Options and the benefits evidenced by this Agreement do not create any entitlement to have the Options or any such benefits transferred to, or assumed by, another company nor to be exchanged, cashed out or substituted for, in connection with any corporate transaction affecting the shares of the Company; and

(15) the following provisions apply only if you are providing services outside the United States:

(i) for employment law purposes outside the United States, the Option, underlying Shares, and the income from and value of same, are not part of normal or expected compensation or salary for any purpose, including but not limited to for purposes of calculating any severance, resignation, termination, redundancy, dismissal, end of service payments, bonuses, holiday pay, long-service awards, pension or retirement benefits or similar mandatory payments; and

(ii) neither the Company, your Employer nor any Affiliate of the Company shall be liable for any foreign exchange rate fluctuation between your local currency and the United States Dollar that may affect the value of the Option or of any amounts due to you pursuant to the exercise of the Option or the subsequent sale of any Shares acquired upon exercise of the Option.

(B) The Company is not providing any tax, legal or financial advice, nor is the Company making any recommendations regarding your participation in the Plan, or your acquisition or sale of the underlying Shares. You should consult with your own personal tax, legal and financial advisors regarding your participation in the Plan before taking any action related to the Plan.

XIII. If one or more of the provisions of this Option shall be held invalid, illegal or unenforceable in any respect, the validity, legality and enforceability of the remaining provisions shall not in any way be affected or impaired thereby and the invalid, illegal or unenforceable provisions shall be deemed null and void; however, to the extent permissible by law, any provisions which could be deemed null and void shall first be construed, interpreted or revised retroactively to permit this Option to be construed so as to foster the intent of this Option and the Plan.

XIV. By electing to accept this Agreement, you acknowledge that you are sufficiently proficient in English or have consulted with an advisor who is sufficiently proficient in English, so as to allow you to understand the terms and conditions of this Agreement. Furthermore, if you have received this Option or any other document related to the Plan translated into a language other than English and if the meaning of the translated version is different than the English version, the English version will control, unless otherwise required by applicable law.

XV. This Option is not intended to constitute “nonqualified deferred compensation” within the meaning of Code Section 409A but rather is intended to be exempt from the application of Code Section 409A. To the extent that this Option is nevertheless deemed to be subject to Code Section 409A for any reason, this Option shall be interpreted in accordance with Code Section 409A and U.S. Department of Treasury regulations and other interpretive guidance issued thereunder, including without limitation any such regulations or other guidance that may be issued after the Grant Date. Notwithstanding any provision herein to the contrary, in the event that following the Grant Date, the Committee (as defined in the Plan) determines that this Option may be or become subject to Code Section 409A, the Committee may adopt such amendments to the Plan and/or this Option or adopt other policies and procedures (including amendments, policies and procedures with retroactive effect), or take any other actions, that the Committee determines are necessary or appropriate to (a) exempt the Plan and/or this Option from the application of Code Section 409A and/or preserve the intended tax treatment of the benefits provided with respect to this Option, or (b) comply with the requirements of Code Section 409A; provided, however, that this paragraph shall not create an obligation on the part of the Committee to adopt any such amendment, policy or procedure or take any such other action.

XVI. By electing to accept this Option, you acknowledge receipt of this Option and hereby confirm your understanding that the terms set forth in this Option constitute, subject to the terms of the Plan, which terms shall control in the event of any conflict between the Plan and this Option, the entire agreement and understanding of the parties with respect to the matters contained herein and supersede any and all prior agreements, arrangements and understandings, both oral and written, between the parties concerning the subject matter of this Option. The Company may, in its sole discretion, decide to deliver any documents related to current or future participation in the Plan (including this Agreement) by electronic means. You hereby consent to receive such documents by electronic delivery and agree to participate in the Plan through an on-line or electronic system established and maintained by the Company or another third party designated by the Company.

XVII. The Company reserves the right to impose other requirements on your participation in the Plan, on this Option and on any Shares acquired under the Plan, to the extent the Company

determines it is necessary or advisable for legal or administrative reasons, and to require you to sign any additional agreements or undertakings that may be necessary to accomplish the foregoing.

XVIII. This Option, the Shares issuable under this Option, and all compensation payable with respect to it shall be subject to clawback, recoupment and/or recovery by the Company pursuant to any and all of the Company's policies with respect to the clawback, recoupment or recovery of compensation in effect as of the Grant Date or as may be adopted or maintained by the Company following the Grant Date, including, without limitation, the Company's Policy on Recovery of Erroneously Awarded Compensation (effective October 2, 2023) and Executive Officer Equity Recoupment Policy (effective December 31, 2020), as they shall be in effect and may be amended from time to time (collectively, the "Clawback Policy"), to the maximum extent permitted by applicable law.

XIX. You acknowledge that a waiver by the Company of breach of any provision of this Option shall not operate or be construed as a waiver of any other provision of this Option, or of any subsequent breach by you or any other grantee.

Very truly yours,

AMGEN INC.

By _____
Duly authorized on behalf
of the Board of Directors

APPENDIX A

ADDITIONAL TERMS AND CONDITIONS OF THE AMGEN INC. SECOND AMENDED AND RESTATED 2009 EQUITY INCENTIVE PLAN, AS AMENDED AND/OR RESTATED FROM TIME TO TIME

GRANT OF STOCK OPTION (BY COUNTRY)

Certain capitalized terms used but not defined in this Appendix A shall have the meanings set forth in the Plan and/or the Agreement to which this Appendix is attached.

TERMS AND CONDITIONS

This Appendix includes additional terms and conditions that govern any Options granted under the Plan if, under applicable law, you are a resident of, are deemed to be a resident of or are working in one of the countries listed below. Furthermore, the additional terms and conditions that govern any Options granted hereunder may apply to you if you transfer employment and/or residency to one of the countries listed below and the Company shall, in its discretion, determine to what extent the terms and conditions contained herein shall apply to you.

NOTIFICATIONS

This Appendix also includes notifications relating to exchange control and other issues of which you should be aware with respect to your participation in the Plan. The information is based on the exchange control, securities and other laws in effect in the countries to which this Appendix refers as of October 2025. Such laws are often complex and change frequently. As a result, the Company strongly recommends that you not rely on the notifications herein as the only source of information relating to the consequences of your participation in the Plan because the information may be outdated when you exercise the Options and acquire Shares under the Plan, or when you subsequently sell Shares acquired under the Plan.

In addition, the notifications are general in nature and may not apply to your particular situation, and the Company is not in a position to assure you of any particular result. Accordingly, you should seek appropriate professional advice as to how the relevant laws in your country may apply to your situation. Finally, if you are a citizen or resident of a country other than the one in which you are currently residing and/or working or are considered a resident of another country for local law purposes, the information contained herein may not be applicable to you or you may be subject to the provisions of one or more jurisdictions.

ALL NON-U.S. JURISDICTIONS

TERMS AND CONDITIONS

Method of Exercise. The following provision replaces Section II(3):

To the extent permitted by applicable statutes and regulations, payment of the exercise price per Share is due in full in cash or check upon exercise of all or any part of this Option which has become exercisable by you. Due to legal restrictions outside the U.S., you are not permitted to pay the exercise price by delivery of already-owned Shares of a value equal to the exercise price of the Shares for which this Option is being exercised. Furthermore, payment may not be made by a combination of cash and already-owned Common Stock.

Tax Withholding. The following provision supplements Section V(C) of the Agreement:

In the event the Company withholds or accounts for Tax Obligations by considering maximum applicable rates in your jurisdiction(s), in the event of over-withholding, you may receive a refund of any over-withheld amount in cash and will not be entitled to the equivalent amount in Shares, or if not refunded, you may seek a refund from the local tax authorities. In the event of under-withholding, you may be required to pay any additional Tax Obligations directly to the applicable tax authority or to the Company and/or your Employer.

NOTIFICATIONS

Insider Trading Restrictions/Market Abuse Laws. You may be subject to insider trading restrictions and/or market abuse laws based on the exchange on which the Shares are listed and in applicable jurisdictions including the United States and your country or your broker's country, if different, which may affect your ability to accept, acquire, sell or otherwise dispose of Shares, rights to Shares (*e.g.*, Options) or rights linked to the value of Shares during such times as you are considered to have "inside information" regarding the Company (as defined by the laws in applicable jurisdictions). Local insider trading laws and regulations may prohibit the cancellation or amendment of orders you place before you possessed inside information. Furthermore, you could be prohibited from (i) disclosing the inside information to any third party, which may include fellow employees (other than on a "need to know" basis) and (ii) "tipping" third parties or causing them otherwise to buy or sell securities. Any restrictions under these laws or regulations are separate from and in addition to any restrictions that may be imposed under any applicable Company insider trading policy. You are responsible for ensuring your compliance with any applicable restrictions and you should speak with your personal legal advisor on this matter.

Foreign Asset/Account, Tax Reporting Information. Your country of residence may have certain foreign asset and/or account reporting requirements which may affect your ability to acquire or hold Shares under the Plan or cash received from participating in the Plan (including from any dividends received, or sale proceeds arising from the sale of Shares) in a brokerage or bank account outside of your country. You may be required to report such accounts, assets or transactions to the tax or other authorities in your country. You also may be required to repatriate sale proceeds or other funds

received as a result of participating in the Plan to your country within a certain time after receipt. You are responsible for ensuring your compliance with such regulations, and you should speak with your personal legal advisor on this matter.

ALL EUROPEAN ECONOMIC AREA (“EEA”) / EUROPEAN UNION (“EU”) JURISDICTIONS, UNITED KINGDOM AND SWITZERLAND

TERMS AND CONDITIONS

Data Privacy Notice. This provision replaces Section IX of the Agreement:

Please refer to the Fair Processing Notice previously provided by your local human resources representative, which notice governs the collection, use and transfer of your personal data necessary for the Company to facilitate your participation in the Plan. If you have any questions or concerns regarding the Fair Processing Notice, including questions about your rights afforded thereunder, you should contact your local human resources representative or send an email to hrconnect@amgen.com.

For purposes of implementing, administering and managing the Plan, Company and your Employer may hold certain personal data about you, including your name, home address and telephone number, email address, date of birth, social insurance/security number (to the extent permitted under applicable local law), passport or other identification number, salary, nationality, job title/work history/service periods, residency status, citizenship, tax withholding and payroll data, any shares of stock or directorships held in the Company, details of all equity compensation or any other entitlement to Shares awarded, cancelled, vested, unvested or outstanding in your favor (“personal data”).

You authorize the transfer of your personal data to Merrill Lynch Bank & Trust Co., FSB, or any successor thereto, and any other third parties which may assist the Company (presently or in the future) with implementing, administering and managing your participation in the Plan to receive, possess, use, retain and transfer your personal data, in electronic or other form, for the purpose of implementing, administering and managing your participation in the Plan, including any requisite transfer of such personal data as may be required to any other broker, escrow agent or other third party with whom the Shares received upon exercise of this Option may be deposited.

ARGENTINA

TERMS AND CONDITIONS

Method of Exercise. Due to legal restrictions in Argentina, you may be required to pay the exercise price for any Shares subject to the Option granted hereunder by a cashless sell-all exercise, such that all Shares will be sold immediately upon exercise and the cash proceeds of sale, less the exercise price, any Tax Obligations and broker’s fees or commissions, will be remitted to you. The Company reserves the right to provide additional methods of exercise depending on local developments.

Labor Law Acknowledgement. The following provision supplements Section XII of the Agreement:

In accepting this Option, you acknowledge, understand and agree that the grant of the Option is made by the Company (not your Employer) in its sole discretion and that the value of the Option or any Shares acquired under the Plan shall not constitute salary or wages for any purpose under Argentine labor law including, but not limited to, the calculation of (i) any labor benefits including, without limitation, vacation pay, thirteenth salary, compensation in lieu of notice, annual bonus, disability, and leave of absence payments, etc., or (ii) any termination or severance indemnities or similar payments.

NOTIFICATIONS

Securities Law Information. The Option and the underlying Shares have not been and will not be publicly issued, placed, distributed, offered or listed in the Argentine capital markets, and, as a result, have not been and will not be registered with the Argentina Securities Commission (Comisión Nacional de Valores, “CNV”). Neither this nor any other offering material related to the Options nor the underlying Shares may be utilized in connection with any general offering to the public in Argentina. Any Argentine resident who acquires Shares under the Plan does so under their own responsibility under the terms of a private offering to the Argentine resident from outside Argentina. Any Argentine resident who acquires Shares shall not transfer such Shares to any other person within six (6) months of acquiring the Shares, unless the transaction is conducted outside Argentina.

Exchange Control Information. Provided you are not required to purchase foreign currency and remit funds out of Argentina to acquire Shares under the Plan, local exchange control restrictions would not apply. However, if so required, you personally are responsible for complying with any and all Argentine currency exchange regulations, approvals and reporting requirements. Exchange control requirements in Argentina are subject to change; you should consult with your personal advisor regarding any obligations you have under the Plan.

Foreign Asset/Account Reporting Information. If you are an Argentine resident, you are required to report certain information regarding any Shares you hold as of December 31 each year to the Argentine tax authorities on your annual tax return.

AUSTRALIA

NOTIFICATIONS

Securities Law Information. If you acquire Shares under the Plan and offer the Shares for sale to a person or entity resident in Australia, the offer may be subject to disclosure requirements under Australian law. You should consult with your own legal advisor before making any such offer in Australia.

Tax Information. Subdivision 83A-C of the Income Tax Assessment Act 1997 (Cth) applies to the Options granted under the Plan, such that the Options are intended to be subject to deferred taxation.

Exchange Control Information. If you are an Australian resident, exchange control reporting is required for cash transactions exceeding a certain threshold (currently AUD10,000) and for international fund transfers. If an Australian bank is assisting with the transaction, the bank will file the report on your behalf. If there is no Australian bank involved in the transfer, you will be required to file the report.

AUSTRIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If you are an Austrian resident and you hold Shares acquired under the Plan outside of Austria, you may be subject to reporting obligations to the Austrian National Bank.

Exchange Control Information.

If the value of Shares meets or exceeds a certain threshold (currently, €5,000,000), you must report the securities held on a quarterly basis to the Austrian National Bank as of the last day of the quarter, on or before the 15th day of the month following the end of the calendar quarter. Further, if you hold cash in accounts outside Austria, monthly reporting requirements will apply, as explained in the next paragraph, if the aggregate transaction volume of such cash accounts meets or exceeds a certain threshold.

A separate reporting requirement applies when you sell Shares acquired under the Plan or receive a cash dividend paid on such Shares. In that case, there may be exchange control obligations if the cash proceeds are held outside of Austria. If the transaction volume of all cash accounts abroad meets or exceeds a certain threshold (currently €10,000,000), the movements and balances of all accounts must be reported monthly, as of the last day of the month, on or before the 15th day of the following month, on the prescribed forms.

BELGIUM

NOTIFICATIONS

Taxation of the Option. Your tax consequences will vary depending on when you accept the Option. If you accept the Option in writing within 60 days of the offer date, you will be subject to taxation on the 60th day after the offer date. If you accept the Option more than 60 days after the offer date, you will be subject to taxation at exercise. Please refer to the additional materials that will be delivered to you for a more detailed description of the tax consequences of accepting the Option. You should consult your personal tax advisor prior to accepting the Option.

Tax Reporting; Foreign Asset/Account Reporting Information. If you are a Belgian resident, you are required to report any securities (*e.g.*, Shares acquired under the Plan) held and bank

accounts (including brokerage accounts) opened and maintained outside of Belgium on your annual tax return. The first time you report the foreign security and/or bank account on your annual income tax return you will have to provide the National Bank of Belgium Central Contact Point with the account details of any such foreign accounts (including the account number, bank name and country in which such account was opened) in a separate form. This report, as well as information on how to complete it, can be found on the website of the National Bank of Belgium, www.nbb.be, under the *Kredietcentrales / Centrales des crédits* caption.

Stock Exchange Tax Information. A stock exchange tax applies to transactions executed by a Belgian resident through a non-Belgian financial intermediary, such as a U.S. broker. The stock exchange tax likely will apply when the Option is exercised and when Shares acquired under the Plan are sold. It is your responsibility to comply with this tax obligation, and you should consult your personal tax advisor for additional details on your obligations with respect to the stock exchange tax.

Annual Securities Accounts Tax Information. An annual securities accounts tax may be payable if the total value of securities held in a Belgian or foreign securities account (e.g., Shares acquired under the Plan) exceeds a certain threshold on four reference dates within the relevant reporting period (i.e., December 31, March 31, June 30 and September 30). In such case, the tax will be due on the value of the qualifying securities held in such account. It is your responsibility to comply with this obligation, and you should consult with your personal tax or financial advisor for additional details.

BRAZIL

TERMS AND CONDITIONS

Compliance with Law. By accepting the Option, you acknowledge that you agree to comply with applicable Brazilian laws and pay any and all applicable taxes associated with the exercise of the Option, the sale of Shares acquired under the Plan and the payment of dividends on such Shares.

Nature of Grant. This provision supplements Section XII of the Agreement:

In accepting this Option, you acknowledge (i) that you are making an investment decision, (ii) that the Options will be exercisable by you only if the vesting conditions are met and any necessary services are rendered by you during the vesting period set forth in the Vesting Schedule, and (iii) that the value of the underlying Shares is not fixed and may increase or decrease in value over the vesting period without compensation to you.

NOTIFICATIONS

Exchange Control Information. If you are resident or domiciled in Brazil, you will be required to submit annually a declaration of assets and rights held outside of Brazil to the Central Bank of Brazil if the aggregate value of such assets and rights on December 31 of each year exceeds US\$1,000,000. If such amount exceeds US\$100,000,000, the referenced declaration must be submitted quarterly, in the month following the end of each quarter. Assets and rights that must be reported include the following: (i) bank deposits; (ii) loans; (iii) financing transactions; (iv)

leases; (v) direct investments; (vi) portfolio investments, including Shares acquired under the Plan; (vii) financial derivatives investments; and (viii) other investments, such as real estate. Please note that foreign individuals holding Brazilian visas are considered Brazilian residents for purposes of this reporting requirement and must declare at least the assets held abroad that were acquired subsequent to the date of admittance as a resident of Brazil. Individuals holding assets and rights outside of Brazil valued at less than US\$1,000,000 are not required to submit a declaration.

Tax on Financial Transactions. Payments to foreign countries, repatriation of funds into Brazil, and the conversion associated with such fund transfers into local currency, may be subject to the Tax on Financial Transactions. It is your responsibility to comply with any applicable Tax on Financial Transactions arising from participation in the Plan and you should consult your personal tax advisor for additional details.

BULGARIA

Exchange Control Information. You will be required to file statistical forms with the Bulgarian National Bank annually regarding your receivables in bank accounts abroad as well as securities held abroad (e.g., Shares acquired under the Plan) if the total sum of all such receivables and securities equals or exceeds a certain threshold as of the previous calendar year-end. The reports are due by March 31. You should contact your bank in Bulgaria for additional information regarding these requirements.

CANADA

TERMS AND CONDITIONS

Termination of Employment. Section IV(B)(1) of the Agreement is amended to read as follows:

- (1) “termination of your employment” shall mean the last date you are either actually providing services to Company or an Affiliate or actually providing services as Director to the Company or an Affiliate; in the event of involuntary termination of your employment (regardless of the reason for such termination and whether or not later found to be invalid or unlawful, including for breaching applicable laws in the jurisdiction where you are employed or the terms of your employment agreement, if any) Except as explicitly and minimally required under applicable legislation, your right, if any, to earn, seek damages in lieu of, vest in, exercise, or otherwise benefit from or participate in any portion of the Option or the Plan will be measured by and immediately terminate, as of the termination of your employment (the “Termination Date”). Except as explicitly and minimally required under applicable legislation, the Termination Date will exclude and will not be extended by any period during which notice, pay in lieu of notice or related payments or damages are provided or required to be provided under applicable laws (including, but not limited to, statute, contract, common/civil law or otherwise). You will not earn or be entitled to any pro-rated vesting or extended exercisability or other benefits under or participation in the Plan for that portion of time before the Termination Date, nor will you be entitled to any compensation for lost vesting, exercisability, benefits, or other participation.

Subject to applicable legislation, if the date you are no longer actually providing services cannot be reasonably determined under the terms of this Agreement or the Plan, the Administrator shall have the exclusive discretion to determine when you are no longer actually providing services for purposes of this Agreement (including whether you may still be considered to be providing services while on a leave of absence). Notwithstanding the foregoing, if applicable employment standards legislation explicitly requires continued vesting or other participation during a statutory notice period, your right to vest in the Options or otherwise participate in or benefit from the Options under the Plan, if any, will terminate effective as of the last day of your minimum statutory notice period, but you will not earn or be entitled to pro-rated vesting or extended exercisability if the vesting date or exercisability period falls after the end of your statutory notice period, nor will you be entitled to any compensation for lost vesting, exercisability, benefits, or other participation;

Nature of Grant. Sections XII(3), XII(12) and XII(15)(i) of the Agreement are amended to read as follows:

(3) no Affiliate (including, but not limited to, the Employer) has any obligation to make any payment of any kind to you under this Agreement, except to the extent explicitly and minimally required under applicable legislation;

(12) in consideration of the grant of this Option, no claim or entitlement to compensation or damages arises from forfeiture of options resulting from (i) the termination of your employment by the Company or an Affiliate (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any) and/or (ii) the application of the Clawback Policy as described in Section XVIII below, except to the extent explicitly and minimally required under applicable legislation;

(15)(i) for employment law purposes outside the United States, the Option, underlying Shares, and the income from and value of same, are not part of normal or expected compensation or salary for any purpose, including but not limited to for purposes of calculating any severance, resignation, termination, redundancy, dismissal, end of service payments, bonuses, holiday pay, long-service awards, pension or retirement benefits or similar mandatory payments, except to the extent explicitly and minimally required under applicable legislation; and

The following provision will apply to you if you are a resident of Quebec:

French Language Documents. A French translation of this document and certain other documents related to this Award will be made available to you as soon as reasonably practicable. You understand that, from time to time, additional information related to the Award may be provided in English and such information may not be immediately available in French. However, upon request, the Company will provide a translation of such information into French as soon as reasonably practicable. Notwithstanding anything to the contrary in the Agreement, and unless you indicate otherwise, the French translation of this document and certain other documents related to the Award will govern your participation in the Plan.

Data Privacy Notice. This provision supplements Section IX of the Agreement:

You hereby authorize the Company and the Company's representative to discuss with and obtain all relevant information from all personnel (professional or not) involved in the administration of the Plan. You further authorize the Company, your Employer and Merrill Lynch Bank & Trust Co., FSB (or any other stock plan service provider) to disclose and discuss your participation in the Plan with their advisors. You also authorize the Company and your Employer to record such information and keep it in your file.

NOTIFICATIONS

Securities Law Information. You are permitted to sell Shares acquired through the Plan through the designated broker appointed under the Plan, if any, provided that the resale of such Shares takes place outside of Canada through the facilities of a stock exchange on which the Shares are listed (e.g., the Nasdaq Global Select Market).

Foreign Asset/Account Reporting Information. Specified foreign property, including Shares, Options and other rights to receive Shares of a non-Canadian company held by a Canadian resident employee generally must be reported annually on a Form T1135 (Foreign Income Verification Statement) if the total cost of the employee's specified foreign property exceeds a certain threshold (currently C\$100,000) at any time during the year. Thus, such Options must be reported – generally at nil cost – if the applicable cost threshold is exceeded because other specified foreign property is held by the employee. When Shares are acquired, their cost generally is the adjusted cost base ("ACB") of the Shares. The ACB ordinarily would equal the fair market value of the Shares at the time of acquisition, but if the employee owns other shares of the same company, this ACB may have to be averaged with the ACB of the other shares.

CHINA

TERMS AND CONDITIONS

The following terms apply only to nationals of the People's Republic of China (the "PRC") residing in the PRC:

Method of Exercise. Due to legal restrictions in the PRC, you will be required to pay the exercise price for any Shares subject to the Option granted hereunder by a cashless sell-all exercise, such that all Shares will be sold immediately upon exercise and the cash proceeds of sale, less the exercise price, any Tax Obligations and broker's fees or commissions, will be remitted to you. The Company reserves the right to provide additional methods of exercise depending on local developments.

Termination of Employment. To comply with requirements imposed by the State Administration of Foreign Exchange, to the extent that, under Section IV of the Agreement, you may exercise any Option after termination of your employment, you will be permitted to exercise such Option for the shorter of the period set forth in Section IV of the Agreement and six (6) months from the date of

termination of your employment; any unexercised Option shall immediately lapse six (6) months following the termination of your employment.

The Company reserves the right to impose such further restrictions or conditions as may be necessary to comply with changes in applicable local laws in the PRC.

Please note that the above provisions will apply to all Options granted to you under the Plan, as well as to any Options granted to you in the past under the Plan.

Exchange Control Requirements. You understand and agree that, pursuant to PRC exchange control requirements, you will be required to repatriate the cash proceeds from the sale of the Shares issued upon the exercise of the Option to China. You further understand that, under applicable laws, such repatriation of your cash proceeds will need to be effectuated through a special exchange control account established by the Company or any Affiliate, including your Employer, and you hereby consent and agree that any proceeds from the sale of the Shares may be transferred to such special account prior to being delivered to you. You also understand that the Company will deliver the proceeds to you as soon as possible, but that there may be delays in distributing the funds to you due to exchange control requirements in China. Proceeds may be paid to you in U.S. dollars or local currency at the Company's discretion. If the proceeds are paid to you in U.S. dollars, you will be required to set up a U.S. dollar bank account in China so that the proceeds may be deposited into this account. If the proceeds are paid to you in local currency, the Company is under no obligation to secure any particular currency conversion rate and the Company may face delays in converting the proceeds to local currency due to exchange control restrictions. You agree to bear any currency fluctuation risk between the date the Option is exercised and the time that (i) the Tax Obligations are converted to local currency and remitted to the tax authorities, and (ii) net proceeds are converted to local currency and distributed to you. You acknowledge that neither the Company nor any Affiliate will be held liable for any delay in delivering the proceeds to you. You agree to sign any agreements, forms and/or consents that may be requested by the Company or the Company's designated broker to effectuate any of the remittances, transfers, conversions or other processes affecting the proceeds. You further agree to comply with any other requirements that may be imposed by the Company in the future in order to facilitate compliance with exchange control requirements in China.

COLOMBIA

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section XII of the Agreement:

You acknowledge that pursuant to Article 15 of Law 50/1990 (Article 128 of the Colombian Labor Code), the Plan and related benefits do not constitute a component of "salary" for any purpose. Therefore, they are considered to be of an extraordinary nature and will not be included and/or considered for purposes of calculating any and all labor benefits, such as legal/fringe benefits, vacations, indemnities, payroll taxes, social insurance contributions and/or any other labor-related amounts, subject to the limitations provided in Law 1393/2010.

Mandate Letter. In accepting the Option, you agree that – if requested by the Company or the Employer – you will execute a Mandate Letter or such other document (whether electronically or by such other method as requested by the Company or the Employer) that the Company determines is necessary or advisable in order that (i) a sufficient number of Shares to be allocated to you upon vesting can be sold on your behalf to cover Tax Obligations required to be withheld by the Employer and (ii) the proceeds from such sale can be wired directly from the Company to the Employer in Colombia for remittance to the tax authorities.

NOTIFICATIONS

Securities Law Information. The Shares are not and will not be registered with the Colombian registry of publicly traded securities (*Registro Nacional de Valores y Emisores*) and therefore the Shares may not be offered to the public in Colombia. Nothing in this document should be construed as the making of a public offer of securities in Colombia.

Exchange Control Information. Investment in assets located abroad (such as Shares acquired under the Plan) does not require prior approval from the Central Bank (*Banco de la República*). Nonetheless, such investments are subject to registration before the Central Bank as foreign investments held abroad, regardless of value.

All payments for your investment originating in Colombia (and the liquidation of such investments) must be transferred through the Colombian foreign exchange market (*e.g.*, local banks), which includes the obligation to correctly complete and file the appropriate foreign exchange form (*declaración de cambio*).

Foreign Asset/Account Reporting Information. An annual information return may need to be filed with the Colombian Tax Office detailing any assets held abroad (including Shares acquired under the Plan). If the individual value of any of these assets exceeds a certain threshold, each asset must be described (*e.g.*, its nature and its value) and the jurisdiction in which it is located must be disclosed. It is your responsibility to comply with this tax reporting requirement.

CROATIA

NOTIFICATIONS

Exchange Control Information. Croatian residents may be required to report any foreign investments (including Shares acquired under the Plan) to the Croatian National Bank for statistical purposes. You should be aware that exchange control regulations in Croatia are subject to frequent change and you are solely responsible for ensuring your continued compliance with current Croatian exchange control laws.

CZECH REPUBLIC

NOTIFICATIONS

Exchange Control Information. If you are a resident of the Czech Republic, you may be required to notify the Czech National Bank (“CNB”) of the acquisition of Shares under the Plan or maintenance of a foreign account if (i) you maintain foreign direct investments with a value of 2,500,000 Kč or more in the aggregate, (ii) you maintain a certain threshold of foreign financial assets, or (iii) you are specifically requested to do so by the CNB.

DENMARK

TERMS AND CONDITIONS

Danish Stock Option Act. In accepting this Option, you acknowledge that you have received an Employer Statement translated into Danish, which is being provided to comply with the Danish Stock Option Act (the “Act”). To the extent more favorable to you and required to comply with the Stock Option Act, as amended with effect from January 1, 2019. You also acknowledge any grant of the Option under the Plan made on or after January 1, 2019 is subject to the rules of the amended Act. Accordingly, you agree that the treatment of the Option upon the termination of your employment is governed solely by Section IV(B)(1) of the Agreement and any corresponding provisions in the Plan. The relevant termination provisions are also detailed in the Employer Statement

Please be aware that as set forth in Section 1 of the Act, the Act only applies to “employees” as that term is defined in Section 2 of the Act. If you are a member of the registered management of an Affiliate or Subsidiary in Denmark or otherwise do not satisfy the definition of employee, you will not be subject to the Act, and the Employer Statement will not apply to you.

NOTIFICATIONS

Foreign Asset/Account Reporting Information. The requirement to report certain information to the Danish Tax Administration via Form V or K was eliminated effective January 1, 2019. However, you still must report the foreign bank/brokerage accounts and their deposits, and Shares held in a foreign bank or brokerage account in your tax return under the section on foreign affairs and income.

EGYPT

NOTIFICATIONS

Exchange Control Information. If you transfer funds into or out of Egypt in connection with the exercise of the Option or the receipt of sale proceeds, you may be required to transfer the funds through a registered bank in Egypt.

FINLAND

NOTIFICATIONS

Foreign Asset/Account Reporting Information. There are no specific reporting requirements with respect to foreign assets/accounts. However, please note that you must check your pre-completed tax return to confirm that the ownership of Shares and other securities (foreign or domestic) are correctly reported. If you find any errors or omissions, you must make the necessary corrections electronically or by sending specific paper forms to the local tax authorities.

FRANCE

TERMS AND CONDITIONS

Language Consent. By accepting the grant, you confirm having read and understood the Plan and Agreement which were provided in the English language. You accept the terms of these documents accordingly.

Consentement Relatif à la Langue Utilisée. *En acceptant l'attribution, vous confirmez avoir lu et compris le Plan et le Contrat, qui ont été communiqués en langue anglaise. Vous acceptez les termes de ces documents en connaissance de cause.*

NOTIFICATIONS

Foreign Asset/Account Reporting Information. French residents and non-residents must declare to the Customs Authorities the cash and securities they import or export without the use of a financial institution when the value of such cash or securities exceeds a certain threshold (currently €10,000). French residents also must report all foreign bank and brokerage accounts on an annual basis (including accounts opened or closed during the tax year) on Form N° 3916, together with the income tax return. Failure to comply could trigger significant penalties.

GERMANY

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If your acquisition of Shares under the Plan leads to a qualified participation at any point during the calendar year, you will need to report the acquisition when you file your tax return for the relevant year. A qualified participation is attained only in the unlikely event (i) you own at least 1% of the Company and the value of the Shares acquired exceeds a certain threshold (currently €150,000) or (ii) you hold Shares exceeding 10% of the Company's total Common Stock.

Exchange Control Information. Cross-border payments in excess of a certain threshold (currently €50,000) must be reported monthly to the German Federal Bank (*Bundesbank*). In case of payments in connection with securities (including proceeds realized upon the sale of Shares or the receipt of dividends), and/or if the Company withholds Shares with a value in excess of the applicable threshold for any Tax Obligations, the report must be made monthly following the month in which the payment was received and must be filed electronically. The form of report (*Allgemeines Meldeportal Statistik*) can be accessed via the *Bundesbank's* website (www.bundesbank.de) and is available in both German

and English. In addition, you may be required to report the acquisition or sale of Shares to the *Bundesbank* if the value of the Shares acquired or sold exceeds the applicable threshold. You are responsible for satisfying any applicable reporting obligation.

GREECE

NOTIFICATIONS

Foreign Asset/Account Reporting Information. The reporting of foreign assets (including Shares and other investments) is your own obligation and takes place through your annual tax return.

HONG KONG

TERMS AND CONDITIONS

Sale of Shares. Shares received at exercise are accepted as a personal investment. In the event that Shares are issued in respect of the Options within six (6) months of the Grant Date, you agree that you will not offer to the public or otherwise dispose of the Shares prior to the six (6)-month anniversary of the Grant Date.

NOTIFICATIONS

SECURITIES WARNING: *The contents of this document have not been reviewed by any regulatory authority in Hong Kong. You should exercise caution in relation to the offer. If you are in doubt about any of the contents of the Agreement, including this Appendix, or the Plan, you should obtain independent professional advice. The Option and any Shares issued in respect of the Option do not constitute a public offering of securities under Hong Kong law and are available only to members of the Board and Employees. The Agreement, including this Appendix, the Plan and other incidental communication materials have not been prepared in accordance with and are not intended to constitute a “prospectus” for a public offering of securities under the applicable securities legislation in Hong Kong. The Option and any documentation related thereto are intended solely for the personal use of each member of the Board and/or Employee and may not be distributed to any other person.*

HUNGARY

There are no country-specific provisions.

ICELAND

TERMS AND CONDITIONS

Method of Exercise. Due to legal restrictions in Iceland, you will be required to pay the exercise price for any Shares subject to the Option granted hereunder by a cashless sell-all exercise, such that all Shares will be sold immediately upon exercise and the cash proceeds of sale, less the exercise price,

any Tax Obligations and broker's fees or commissions, will be remitted to you. The Company reserves the right to provide additional methods of exercise depending on local developments.

NOTIFICATIONS

Exchange Control Information. Approval by the Central Bank of Iceland is no longer required to participate in the Plan, regardless of the value of the Shares acquired under the Plan. Despite the recent relaxation of the exchange control requirements, you should consult with your personal advisor to ensure compliance with applicable exchange control regulations in Iceland as such regulations are subject to frequent change. You are responsible for ensuring compliance with all exchange control laws in Iceland.

INDIA

TERMS AND CONDITIONS

Method of Exercise. Due to legal restrictions in India, you will not be permitted to pay the exercise price for Shares subject to the Option granted hereunder by a cashless "sell-to-cover" procedure, under which method a number of Shares with a value sufficient to cover the exercise price, brokerage fees and any applicable Tax Obligations would be sold upon exercise and you would receive only the remaining Shares subject to the exercised Option. The Company reserves the right to permit this procedure for payment of the exercise price in the future, depending on the development of local law.

NOTIFICATIONS

Exchange Control Information. If you remit funds from India to pay the exercise price, you may be subject to Tax Collection At Source ("**TCS**") if your annual remittances out of India exceed a certain amount (currently INR 1,000,000). You may be required to provide a declaration to the bank remitting the funds to determine if the TCS limit has been reached. If deemed necessary to comply with applicable laws, the Company may require you to pay for the shares purchased on exercise, and any Tax Obligations through a cashless "sell-all" exercise or net exercise method. The Company reserves the right to prescribe alternative methods of payment depending on the development of local laws.

You must repatriate any funds received in connection with the Option (e.g., proceeds from the Shares and the receipt of dividends) within such time as prescribed under applicable Indian exchange control laws, which may be amended from time to time, unless an exemption applies. You should obtain a foreign inward remittance certificate ("**FIRC**") from the bank in which you deposit the foreign currency and maintain the FIRC as evidence of the repatriation of funds in the event the Reserve Bank of India or the Employer requests proof of repatriation. It is your responsibility to comply with these requirements. Neither the Company nor the Employer will be liable for any fines or penalties resulting from your failure to comply with any applicable laws. You may be required to provide information regarding funds received from participation in the

Plan to the Company and/or the Employer to enable them to comply with their filing requirements under exchange control laws in India.

Foreign Asset/Account Reporting Information. You are required to declare foreign bank accounts and any foreign financial assets (including Shares held outside of India) in your annual tax return. It is your responsibility to comply with this reporting obligation, and you should consult your personal tax advisor in this regard.

IRELAND

TERMS AND CONDITIONS

Nature of Grant. This provision supplements Section XII of the Agreement:

In accepting this Option, you acknowledge that the benefits received under the Plan will not be taken into account for any redundancy or unfair dismissal claim.

ITALY

TERMS AND CONDITIONS

Method of Exercise. Due to legal restrictions in Italy, you will be required to pay the exercise price for any Shares subject to the Option granted hereunder by a cashless sell-all exercise, such that all Shares will be sold immediately upon exercise and the cash proceeds of sale, less the exercise price, any Tax Obligations and broker's fees or commissions, will be remitted to you. The Company reserves the right to provide additional methods of exercise depending on local developments.

Nature of Grant. In accepting this Option, you acknowledge that (1) you have received a copy of the Plan, the Agreement and this Appendix; (2) you have reviewed the applicable documents in their entirety and fully understand the contents thereof; and (3) you accept all provisions of the Plan, the Agreement and this Appendix.

For the Option granted, you further acknowledge that you have read and specifically and explicitly approve, without limitation, the following Sections of the Option Agreement: Section I, Section IV, Section V, Section X, Section XII, Section XIII, Section XIV, Section XVII and the Data Privacy Notice for All European Economic Area ("EEA") / European Union ("EU") Jurisdictions, United Kingdom and Switzerland in this Appendix.

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Italian residents who, at any time during the fiscal year, hold foreign financial assets (including cash and Shares) which may generate income taxable in Italy are required to report these assets on their annual tax returns (UNICO Form, RW Schedule) for the year during which the assets are held, or on a special form if no tax return is due. These reporting

obligations will also apply to Italian residents who are the beneficial owners of foreign financial assets under Italian money laundering provisions.

Foreign Financial Assets Tax. The fair market value of any Shares held outside of Italy is subject to a foreign assets tax at a flat rate. The fair market value is considered to be the value of the Shares on the Nasdaq Global Select Market on December 31 of the applicable year in which you held the Shares (or when the Shares are acquired during the course of the year, the tax is levied in proportion to the actual days of holding over the calendar year). No tax payment duties arise if the amount of the foreign financial assets tax calculated on all financial assets held abroad does not exceed a certain threshold. You should consult with your personal tax advisor about the foreign financial assets tax.

JAPAN

NOTIFICATIONS

Exchange Control Information. If you acquire Shares valued at more than ¥100,000,000 in a single transaction, you must file a Securities Acquisition Report with the Ministry of Finance through the Bank of Japan within 20 days of the purchase of the Shares.

In addition, if you pay more than ¥30,000,000 in a single transaction for the purchase of Shares when you exercise the Option, you must file a Payment Report with the Ministry of Finance through the Bank of Japan by the 20th day of the month following the month in which the payment was made. The precise reporting requirements vary depending on whether or not the relevant payment is made through a bank in Japan.

A Payment Report is required independently from a Securities Acquisition Report. Therefore, if the total amount that you pay upon a one-time transaction for exercising the Option and purchasing Shares exceeds ¥100,000,000, then you must file both a Payment Report and a Securities Acquisition Report.

Foreign Asset/Account Reporting Information. You will be required to report to the Japanese tax authorities details of any assets held outside of Japan as of December 31st (including any Shares acquired under the Plan) to the extent such assets have a total net fair market value exceeding a certain threshold (currently ¥50,000,000). Such report will be due by June 30 each year. You should consult with your personal tax advisor as to whether the reporting obligation applies to you and whether you will be required to include in the report details of any outstanding Options, Shares or cash that you hold.

KOREA (SOUTH)

NOTIFICATIONS

Exchange Control Information. Korean residents who sell Shares acquired under the Plan and/or receive cash Dividends on the Shares must file a report with a Korean foreign exchange bank if the proceeds exceed a certain threshold (currently US\$5,000 per transaction) and are deposited

into a non- Korean bank account. The report is not required if proceeds are deposited into a non-Korean brokerage account. It is your responsibility to ensure compliance with any applicable exchange control reporting obligations.

Foreign Asset/Account Reporting Information. To remit funds out of Korea to exercise the Option by means of a cash exercise method, you must obtain a confirmation of the remittance by a foreign exchange bank in Korea. This is an automatic procedure (*i.e.*, the bank does not need to approve the remittance and the process should not take more than a single day). You likely will need to present to the bank processing the transaction supporting documentation evidencing the nature of the remittance.

You are required to declare all foreign financial accounts (*e.g.* non-Korean bank accounts, brokerage accounts holding Shares, etc.) to the Korean tax authority and file a report regarding such accounts if the monthly balance of such accounts exceeds a certain threshold. It is your responsibility to comply with this reporting obligation, and you should consult your personal tax advisor to ensure compliance with this requirement.

LITHUANIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If you (i) hold certain job positions established by the law or (ii) donate to political parties or political campaigners, you must file an Annual Asset Return of the Individual (Family) in Form No. FR0001 with respect to assets held outside of Lithuania (*e.g.*, Shares). If you open an account in a foreign financial institution and annual turnover in the account exceeds a certain threshold (currently €15,000), you must file a foreign account report.

MEXICO

TERMS AND CONDITIONS

Acknowledgement of the Agreement. In accepting the Option granted hereunder, you acknowledge that you have received a copy of the Plan, have reviewed the Plan and the Option Agreement, including this Appendix, in their entirety and fully understand and accept all provisions of the Plan and the Agreement, including this Appendix. You further acknowledge that you have read and specifically and expressly approve the terms and conditions of Section XII of the Agreement, in which the following is clearly described and established:

- (1) Your participation in the Plan does not constitute an acquired right.
- (2) The Plan and your participation in the Plan are offered by Amgen Inc. on a wholly discretionary basis.
- (3) Your participation in the Plan is voluntary.

- (4) Amgen Inc. and its Affiliates are not responsible for any decrease in the value of the Option granted and/or Shares issued under the Plan.

Labor Law Acknowledgement and Policy Statement. In accepting the Option granted hereunder, you expressly recognize that Amgen Inc., with registered offices at One Amgen Center Drive, Thousand Oaks, California 91320, U.S.A., is solely responsible for the administration of the Plan and that your participation in the Plan and acquisition of Shares do not constitute an employment relationship between you and Amgen Inc. since you are participating in the Plan on a wholly commercial basis and your sole employer is Amgen Mexico S.A. de C.V. ("Amgen-Mexico"). Based on the foregoing, you expressly recognize that the Plan and the benefits that you may derive from participation in the Plan do not establish any rights between you and your Employer, Amgen-Mexico, and do not form part of the employment conditions and/or benefits provided by Amgen-Mexico and any modification of the Plan or its termination shall not constitute a change or impairment of the terms and conditions of your employment.

You further understand that your participation in the Plan is as a result of a unilateral and discretionary decision of Amgen Inc.; therefore, Amgen Inc. reserves the absolute right to amend and/or discontinue your participation in the Plan at any time without any liability to you.

Finally, you hereby declare that you do not reserve to yourself any action or right to bring any claim against Amgen Inc. for any compensation or damages regarding any provision of the Plan or the benefits derived under the Plan, and you therefore grant a full and broad release to Amgen Inc., its Affiliates, stockholders, officers, agents or legal representatives with respect to any claim that may arise.

Spanish Translation

Reconocimiento del Otorgamiento. Al aceptar cualquier Opción bajo el presente documento, usted reconoce que ha recibido una copia del Plan, que ha revisado el mismo en su totalidad, así como también el Acuerdo de Opción, incluyendo este Apéndice, además que comprende y está de acuerdo con todas las disposiciones tanto del Plan y del Opción, incluyendo este Apéndice. Asimismo, usted reconoce que ha leído y manifiesta específicamente y expresamente la conformidad con los términos y condiciones establecidos en la Sección XII del Acuerdo de Opción, en los que se establece y describe claramente que:

- (1) Su participación en el Plan de ninguna manera constituye un derecho adquirido.
- (2) El Plan y su participación en el mismo son ofrecidos por Amgen Inc. de forma completamente discrecional.
- (3) Su participación en el Plan es voluntaria.
- (4) Amgen Inc. y sus Afiliados no son responsables de ninguna disminución en el valor de la opción otorgada y/o de las Acciones Comunes emitidas mediante el Plan.

Reconocimiento de la Ley Laboral y Declaración de Política. Al aceptar cualquier Opción bajo el presente, usted reconoce expresamente que Amgen Inc., con oficinas registradas localizadas en One Amgen Center Drive, Thousand Oaks, California 91320, U.S.A., es la única responsable de la administración del Plan y que su participación en el mismo y la adquisición de Acciones Comunes no constituyen de ninguna manera una relación laboral entre usted y Amgen Inc., debido a que su participación en el Plan es únicamente una relación comercial y que su único empleador es Amgen Mexico S.A. de C.V. (“Amgen-México”). Derivado de lo anterior, usted reconoce expresamente que el Plan y los beneficios a su favor que pudieran derivar de la participación en el mismo, no establecen ningún derecho entre usted y su empleador, Amgen – México, y no forman parte de las condiciones laborales y/o los beneficios otorgados por Amgen – México, y cualquier modificación del Plan o la terminación del mismo no constituirá un cambio o desmejora de los términos y condiciones de su trabajo.

Asimismo, usted entiende que su participación en el Plan es resultado de la decisión unilateral y discrecional de Amgen Inc., por lo tanto, Amgen Inc. se reserva el derecho absoluto de modificar y/o discontinuar su participación en el Plan en cualquier momento y sin ninguna responsabilidad para usted.

Finalmente, usted manifiesta que no se reserva ninguna acción o derecho que origine una demanda en contra de Amgen Inc., por cualquier compensación o daños y perjuicios, en relación con cualquier disposición del Plan o de los beneficios derivados del mismo, y en consecuencia usted exime amplia y completamente a Amgen Inc. de toda responsabilidad, como así también a sus Afiliadas, accionistas, directores, agentes o representantes legales con respecto a cualquier demanda que pudiera surgir.

NOTIFICATIONS

Securities Law Information. The Options and the Shares offered under the Plan have not been registered with the National Register of Securities maintained by the Mexican National Banking and Securities Commission and cannot be offered or sold publicly in Mexico. In addition, the Plan, the Agreement and any other document relating to the Options may not be publicly distributed in Mexico. These materials are addressed to you only because of your existing relationship with the Company and your Employer and these materials should not be reproduced or copied in any form. The offer contained in these materials does not constitute a public offering of securities but rather constitutes a private placement of securities addressed specifically to individuals who are present employees of Amgen-Mexico made in accordance with the provisions of the Mexican Securities Market Law, and any rights under such offering shall not be assigned or transferred.

NETHERLANDS

NOTIFICATIONS

Securities Law Information.

**Attention! This investment falls outside AFM supervision.
No prospectus required for this activity.**



NORWAY

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Norwegian residents may be subject to foreign asset reporting as part of their ordinary tax return. Norwegian banks, financial institutions, limited companies etc. must report certain information to the Tax Administration. Such information may then be pre-completed in a Norwegian resident's tax return. However, if the resident has traded, or is the owner of, financial instruments (*e.g.*, Shares) not pre-completed in the tax return, the Norwegian resident must enter this information in Form RF-1159, which is an appendix to the tax return.

Options will be considered assets and are, therefore, subject to wealth tax. An exemption from wealth tax may be available for non-transferrable awards. However, because the wealth tax regulations and the practice of the tax authorities are not well developed, Norwegian residents should provide the tax authorities with information concerning the Options in the annual tax return even if the Norwegian resident maintains that no wealth tax is payable.

Exchange Control Information. In general, Norwegian residents should not be subject to any foreign exchange requirements in connection with their acquisition or sale of Shares under the Plan, except normal reporting requirements to the Norwegian Currency Registry. If any transfer of funds into or out of Norway is made through a Norwegian bank, the bank will make the registration.

POLAND

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Polish residents holding foreign securities (including Shares) and maintaining accounts abroad must file reports with the National Bank of Poland if the aggregate value of cash and securities held in such foreign accounts exceeds a certain threshold (currently PLN 7 million). If required, the reports are due on a quarterly basis by the 20th day following the end of each quarter and must be filed on special forms available on the website of the National Bank of Poland.

Exchange Control Information. In addition, Polish residents are required to transfer funds through a bank account in Poland if the transferred amount in any single transaction exceeds a specified threshold (currently €15,000 (or PLN 15,000 if such transfer of funds is associated with the business activity of a consultant)). You must store all documents connected with any foreign exchange transactions you engage in for a period of five (5) years from the end of the year when such transactions were made. Penalties may apply for failure to comply with exchange control requirements.

PORTUGAL

TERMS AND CONDITIONS

Consent to Receive Information in English. You hereby expressly declare that you have full knowledge of the English language and have read, understood and fully accepted and agreed with the terms and conditions established in the Plan and Agreement.

Conhecimento da Língua. *Por meio do presente, eu declaro expressamente que tem pleno conhecimento da língua inglesa e que li, compreendi e livremente aceitei e concordei com os termos e condições estabelecidas no Plano e no Acordo.*

PUERTO RICO

There are no country-specific provisions.

ROMANIA

NOTIFICATIONS

Exchange Control Information. Certain transfers of funds may need to be reported to the National Office for Prevention and Control of Money Laundering on specific forms by the relevant bank or financial institution. If you deposit proceeds from the sale of Shares or the receipt of dividends in a bank account in Romania, you may be required to provide the Romanian bank assisting with the transaction with appropriate documentation explaining the source of the income. You should consult with a legal advisor to determine whether you will be required to submit such documentation to the Romanian bank.

RUSSIA

NOTIFICATIONS

Foreign Account Report Information. Russian residents maintaining accounts abroad must disclose such accounts with the Russian tax authorities, as follows: (i) file notifications upon opening (or closing or changing bank details) within 30 days, and (ii) file reports on the movement of cash or financial assets on such accounts by June 1 of the following year. Such notifications and reports must be filed on special forms available on the website of the Russian

tax authorities. Russian residents who are absent from Russia for more than 183 days in a year may be relieved from this reporting.

TERMS AND CONDITIONS

Securities Law Information. The sale of Shares acquired under the Plan is not permitted within Russia. The Shares are currently traded on the Nasdaq Global Select Market in the U.S. under the ticker symbol “AMGN,” therefore, the Shares may be sold on this exchange.

You understand that the exchange control rules and regulations in Russia and the legal restrictions impacting your participation in the Plan are subject to frequent change. You should consult with your personal legal advisor to determine the applicability of any requirements or restrictions applicable to any Shares or cash received in connection with the Plan.

SINGAPORE

TERMS AND CONDITIONS

Restriction on Sale and Transferability. You hereby agree that any Shares acquired pursuant to the Option will not be offered for sale in Singapore prior to the six (6)-month anniversary of the Grant Date, unless such sale or offer is made pursuant to one or more exemptions under Part XIII Division 1 Subdivision (4) (other than section 280) of the Securities and Futures Act (Chap. 289, 2006 Ed.) (“SFA”), or pursuant to, and in accordance with the conditions of, any other applicable provisions of the SFA.

NOTIFICATIONS

Securities Law Information. The grant of the Option is being made pursuant to the “Qualifying Person” exemption under section 273(1)(f) of the SFA, on which basis it is exempt from the prospectus and registration requirements under the SFA, and is not made with a view to the Option being subsequently offered for sale to any other party. The Plan has not been, and will not be, lodged or registered as a prospectus with the Monetary Authority of Singapore.

Director Notification Requirement. Directors (including alternate, substitute, associate and shadow directors) of a Singapore Affiliate are subject to certain notification requirements under the Singapore Companies Act, regardless of whether they are resident or employed in Singapore. Directors of a Singapore Affiliate must notify the Singapore Affiliate in writing of an interest (*e.g.*, Options, Shares, etc.) in the Company or any related company within two (2) business days of (i) its acquisition or disposal, (ii) any change in a previously disclosed interest (*e.g.*, when the Shares are sold), or (iii) becoming a director.

SLOVAK REPUBLIC

There are no country-specific provisions.

SLOVENIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Slovenian residents may be required to report the opening of bank and/or brokerage accounts to tax authorities within eight (8) days of opening such account. You should consult with your personal tax advisor to determine whether this requirement will be applicable to any accounts opened in connection with your participation in the Plan (*e.g.*, your brokerage account with the Company's designated broker).

SPAIN

TERMS AND CONDITIONS

Nature of Grant. The following provision supplements Section XII of the Agreement:

In accepting this Option, you consent to participation in the Plan and acknowledge that you have received a copy of the Plan.

You understand that the Company has unilaterally, gratuitously and in its sole discretion decided to grant the Option under the Plan to individuals who may be members of the Board or Employees of the Company or its Affiliates throughout the world. The decision is a limited decision, which is entered into upon the express assumption and condition that (i) the Option granted will not economically or otherwise bind the Company or any of its Affiliates on an ongoing basis, other than as expressly set forth in the Agreement, including this Appendix; (ii) Option granted hereunder and any Shares acquired under the Plan shall not become a part of any employment contract (either with the Company or any of its Affiliates) and shall not be considered a mandatory benefit, salary for any purposes (including severance compensation) or any other right whatsoever; and (iii) any unvested Options will be forfeited immediately upon termination of your employment for any reason, as detailed below. Further, you understand and freely accept that there is no guarantee that any benefit whatsoever shall arise from any gratuitous and discretionary grant of the Option since the future value of the Option and the underlying Shares is unknown and unpredictable. In addition, you understand that the Option granted hereunder would not be made but for the assumptions and conditions referred to above; thus, you understand, acknowledge and freely accept that, should any or all of the assumptions be mistaken or should any of the conditions not be met for any reason, then any grant of an Option or right to an Option shall be null and void.

Further, the vesting of the Option is expressly conditioned on your continued and active rendering of service, such that if your employment terminates for any reason whatsoever, the Option may cease vesting immediately, in whole or in part, effective on the date of your termination of employment (unless otherwise specifically provided in Section IV of the Agreement). This will be the case, for example, even if (1) you are considered to be unfairly dismissed without good cause (*i.e.*, subject to a “*despido improcedente*”); (2) you are dismissed for disciplinary or objective reasons or due to a collective dismissal; (3) you terminate service due to a change of work location, duties or any other employment or contractual condition; (4) you terminate service due to a unilateral breach of contract by the Company or an Affiliate; or (5) your employment terminates for any other reason whatsoever.

Consequently, upon termination of your employment for any of the above reasons, you may automatically lose any rights to Options that were not vested on the date of your termination of employment, as described in the Plan and the Agreement.

You acknowledge that you have read and specifically accept the conditions referred to in Section IV of the Agreement.

NOTIFICATIONS

Securities Law Information. No “offer of securities to the public,” as defined under Spanish law, has taken place or will take place in the Spanish territory. The Agreement (including this Appendix) has not been nor will it be registered with the *Comisión Nacional del Mercado de Valores*, and does not constitute a public offering prospectus.

Exchange Control Information. You will be required to declare electronically to the Bank of Spain any security accounts (including brokerage accounts held abroad), as well as the securities (including Shares acquired under the Plan) held in such accounts if the value of the transactions for all such accounts during the prior year or the balances of such accounts as of December 31 of the prior year exceeds a certain threshold (currently, €1,000,000). Different thresholds and deadlines to file this declaration apply. However, if neither such transactions during the immediately preceding year nor the balances / positions as of December 31 exceed the applicable threshold, no such declaration must be filed unless expressly required by the Bank of Spain. If any of such thresholds were exceeded during the current year, you may be required to file the relevant declaration corresponding to the prior year, however, a summarized form of declaration may be available..

Foreign Asset/Account Reporting Information.

To the extent that you hold Shares and/or have bank accounts outside of Spain with a value in excess of €50,000 (for each type of asset) as of December 31 each year, you will be required to report information on such assets in your tax return (tax form 720) for such year. After such Shares and/or accounts are initially reported, the reporting obligation will apply for subsequent years only if the value of any previously-reported Shares or accounts increases by more than €20,000 (for each type of asset) as of each subsequent December 31, or if you sell or otherwise dispose of any previously-reported Shares or accounts. If the value of such Shares and/or accounts as of December 31 does not exceed €50,000, a summarized form of declaration may be presented.

SWEDEN

TERMS AND CONDITIONS

Authorization to Withhold. This provision supplements Section V of the Agreement:

Without limiting the Company’s and the Employer’s authority to satisfy their withholding obligations for Tax Obligations as set forth in the Agreement, in accepting the Options, you authorize the Company to withhold Shares or to sell Shares otherwise issuable to you upon exercise to satisfy Tax Obligations, regardless of whether the Company and/or Employer have an obligation to withhold such Tax

Obligations, provided that such withholding would not, in the Company's determination, result in adverse accounting consequences to the Company.

SWITZERLAND

NOTIFICATIONS

Securities Law Information. Neither this document nor any other materials relating to the Option (i) constitutes a prospectus according to articles 35 et seq. of the Swiss Federal Act on Financial Services ("**FinSA**"), (ii) may be publicly distributed or otherwise made publicly available in Switzerland to any person other than an employee of the Company or one of its Subsidiaries or (iii) has been or will be filed with, approved or supervised by any Swiss reviewing body according to article 51 of FinSA or any Swiss regulatory authority, including the Swiss Financial Market Supervisory Authority.

TAIWAN

NOTIFICATIONS

Securities Law Information. Participation in the Plan is available only for select Board members and Employees of the Company and its Affiliates. The grant of Units and participation in the Plan is not a public offer of securities by a Taiwanese company.

Exchange Control Information. You may acquire and remit foreign currency (including proceeds from the sale of Shares or the receipt of dividends) up to US\$10,000,000 per year without justification. If the transaction amount is TWD500,000 or more in a single transaction, you must submit a Foreign Exchange Transaction Form. If the transaction amount is US\$500,000 or more in a single transaction, you must also provide supporting documentation to the satisfaction of the remitting bank.

THAILAND

NOTIFICATIONS

Exchange Control Information. If you receive funds in connection with the Plan (*e.g.*, dividends or sale proceeds) with a value equal to or greater than a certain threshold (currently US\$1,000,000) per transaction, you are required to immediately repatriate such funds to Thailand. Any foreign currency repatriated to Thailand must be converted to Thai Baht or deposited into a foreign currency deposit account opened with any commercial bank in Thailand acting as the authorized agent within 360 days from the date the funds are repatriated to Thailand. You are also required to inform the authorized agent of the details of the foreign currency transaction, including your identification information and the purpose of the transaction. The Employee is responsible for ensuring compliance with all exchange control laws in Thailand.

If you do not comply with the above obligations, you may be subject to penalties assessed by the Bank of Thailand. Because exchange control regulations change frequently and without notice,

you should consult your legal advisor before selling any Shares (or receiving any other funds in connection with the Plan) to ensure compliance with current regulations. It is your responsibility to comply with exchange control laws in Thailand, and neither the Company nor your Employer will be liable for any fines or penalties resulting from failure to comply with applicable laws.

TÜRKIYE

NOTIFICATIONS

Securities Law Information. The sale of Shares acquired under the Plan is not permitted within Türkiye. The sale of Shares acquired under the Plan must occur outside of Türkiye. The Shares are currently traded on the Nasdaq Global Select Market in the U.S. under the ticker symbol “AMGN” and Shares may be sold on this exchange.

Exchange Control Information. You may be required to engage a Turkish financial intermediary to assist with the cash exercise of an Option or the sale of Shares acquired under the Plan. To the extent a Turkish financial intermediary is required in connection with the Option exercise or the sale of any Shares acquired upon exercise of the Option, you are solely responsible for engaging such Turkish financial intermediary. You should consult your personal legal advisor prior to the exercise of Options or any sale of Shares to ensure compliance with the current requirements.

UNITED ARAB EMIRATES

NOTIFICATIONS

Securities Law Information. Options under the Plan are granted only to select Board members and Employees of the Company and its Affiliates and are for the purpose of providing equity incentives. The Plan and the Agreement are intended for distribution only to such Board members and Employees and must not be delivered to, or relied on by, any other person. You should conduct your own due diligence on the Options offered pursuant to this Agreement. If you do not understand the contents of the Plan and/or the Agreement, you should consult an authorized financial adviser. The Emirates Securities and Commodities Authority and the Dubai Financial Services Authority have no responsibility for reviewing or verifying any documents in connection with the Plan. Further, the Ministry of the Economy and the Dubai Department of Economic Development have not approved the Plan or the Agreement nor taken steps to verify the information set out therein, and have no responsibility for such documents.

UNITED KINGDOM

TERMS AND CONDITIONS

Tax Withholding. This provision supplements Section V of the Agreement:

Without limitation to Section V of the Agreement, you agree that you are liable for all Tax Obligations and hereby covenant to pay all such Tax Obligations as and when requested by the Company or your

Employer or by HM Revenue and Customs (“**HMRC**”) (or any other tax authority or any other relevant authority). You also agree to indemnify and keep indemnified the Company and your Employer against any taxes that they are required to pay or withhold or have paid or will pay to HMRC (or any other tax authority or any other relevant authority) on your behalf.

Notwithstanding the foregoing, if you are an executive officer or director within the meaning of Section 13(k) of the Exchange Act, as amended from time to time, you understand that you may not be able to indemnify the Company or your Employer for the amount of income tax not collected from or paid by you, as it may be considered a loan. In the event that you are an executive officer or director and income tax is not collected from you within ninety (90) days after the end of the tax year in which the Taxable Event occurs, the amount of any uncollected income tax may constitute an additional benefit to you on which additional income tax and national insurance contributions (“**NICs**”) may be payable. You acknowledge that you are responsible for reporting and paying any income tax due on this additional benefit directly to HMRC under the self-assessment regime and for paying your Employer for the amount of any NICs due on this additional benefit, which the Company or your Employer may obtain from you by any of the means set forth in Section V of the Agreement.

If the maximum applicable withholding rate is used, any over-withheld amount may be credited to you by the Company or your Employer (with no entitlement to the Common Stock equivalent) or if not so credited, you may seek a refund from the local tax authorities.

Joint Election. If you are a resident of the United Kingdom between the Grant Date and the vesting of the Option, as a condition of the Option granted hereunder, you agree to accept any liability for secondary Class 1 National Insurance Contributions (the “**Employer NICs**”), which may be payable by the Company or your Employer with respect to the exercise of the Option and issuance of Shares subject to the Option, the assignment or release of the Option for consideration, or the receipt of any other benefit in connection with the Option.

Without limitation to the foregoing, you agree to make an election (the “**Election**”), in the form specified and/or approved for such election by HMRC, that the liability for your Employer NICs payments on any such gains shall be transferred to you to the fullest extent permitted by law. You further agree to execute such other elections as may be required between you and any successor to the Company and/or your Employer. You hereby authorize the Company and your Employer to withhold such Employer NICs by any of the means set forth in Section V of the Agreement.

Failure by you to enter into an Election, withdrawal of approval of the Election by HMRC or a joint revocation of the Election by you and the Company or your Employer, as applicable, shall be grounds for the forfeiture and cancellation of the Option, without any liability to the Company or your Employer.

UNITED STATES

TERMS AND CONDITIONS

Nature of Grant. The following provision replaces Section IV(B)(1) of the Agreement:

(1) “termination of your employment” shall mean the last date you are either an active employee of the Company or an Affiliate or actively engaged as a Director of the Company or an Affiliate; in the event of termination of your employment (whether or not in breach of local labor laws), your right to exercise the Option and vest under the Plan, if any, will terminate effective as of the date that you are no longer actively employed; provided, however, that such right will be extended by any notice period mandated by law (e.g. the Worker Adjustment and Retraining Notification Act (“WARN Act”) notice period or similar periods pursuant to local law) and any paid administrative leave (as applicable), unless the Company shall provide you with written notice otherwise before the commencement of such notice period or leave. Your right, if any, to exercise the Option after termination of employment will be measured by the date of termination of your active employment; provided, however, that such right will be extended by any notice period mandated by law (e.g. the Worker Adjustment and Retraining Notification Act (“WARN Act”) notice period or similar periods pursuant to local law) and any paid administrative leave, unless the Company shall provide you with written notice otherwise before the commencement of such notice period or leave. Notwithstanding anything to the contrary herein, in no event shall the term of this Option extend beyond the Expiration Date set forth on the Award Notice and in this Agreement.

Form of Award Notice

[The information set forth in this Award Notice will be contained on the related pages on Merrill Lynch Benefits Website (or the website of any successor company to Merrill Lynch Bank & Trust Co., FSB). This Award Notice shall be replaced by the equivalent pages on such website. References to Award Notice in this Agreement shall then refer to the equivalent pages on such website.]

This notice of Award (the “Award Notice”) sets forth certain details relating to the grant by the Company to you of the Award identified below, pursuant to the Plan. The terms of this Award Notice are incorporated into the Restricted Stock Unit Agreement (the “Agreement”) that accompanies this Award Notice and made part of the Agreement. Capitalized terms used in this Award Notice that are not otherwise defined in this Award Notice have the meanings given to such terms in the Agreement.

Employee:

Employee ID:

Address:

Award Type:

Grant ID:

Plan: Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan, as amended and/or restated from time to time

Grant Date:

Grant Price: \$ _____

Number of Shares:

Number of Units

Vesting Date: Means the vesting date indicated in the Vesting Schedule

Vesting Schedule: Means the schedule of vesting set forth under Vesting Details

Vesting Details: Means the presentation (tabular or otherwise) of the Vesting Date and the quantity of Shares vesting

IMPORTANT NOTICE REGARDING ACCEPTANCE OF THE AWARD AND THE REQUIREMENT TO OPEN A BROKERAGE ACCOUNT¹:

RESIDENTS OF THE U.S. AND PUERTO RICO: Please read this Award Notice, the Plan and the Agreement (collectively, the “Grant Documents”) carefully. If you, as a resident of the U.S. or Puerto Rico, do **not** wish to receive this Award and/or you do **not** consent and agree to the terms and conditions on which this Award is offered, as set forth in the Grant Documents, then you must reject the Award by contacting the Merrill Lynch call center at +1 (800) 97AMGEN (+1 (800) 972-6436) within the U.S., Puerto Rico and Canada or +1 (609) 818-8910 from all other countries (Merrill Lynch will accept the charges for your call) no later than the forty-fifth calendar day following the day on which this Award Notice is made available to you,

¹ This provision is only for use on the form of grant used for the U.S. and Puerto Rico.

in which case the Award will be cancelled. For the purpose of determining the forty-five calendar days, Day 1 will be the day **immediately** following the day on which this Award Notice is made available to you. Your failure to notify the Company of your rejection of the Award or your refusal of, or disagreement with, all terms and conditions of the Award, as set forth in the Grant Documents, within this specified period will constitute your acceptance of the Award and your agreement with all terms and conditions of the Award, as set forth in the Grant Documents. If you agree to the terms and conditions of your grant and you desire to accept it, then no further action is needed on your part to accept the grant. However, you must still open a brokerage account as directed by the Company, by 1:00 pm Pacific Time on or before the date that is 11 months after the Grant Date specified in this Award Notice. This step is necessary to process transactions related to your equity grant. If you do not open a brokerage account by this deadline, **your grant will be cancelled.**

RESTRICTED STOCK UNIT AGREEMENT

THE SPECIFIC TERMS OF YOUR GRANT OF RESTRICTED STOCK UNITS ARE FOUND IN THE PAGES RELATING TO THE GRANT OF RESTRICTED STOCK UNITS FOUND ON MERRILL LYNCH BENEFITS WEBSITE (OR THE WEBSITE OF ANY SUCCESSOR COMPANY TO MERRILL LYNCH BANK & TRUST CO., FSB) (THE “AWARD NOTICE”) WHICH ACCOMPANIES THIS DOCUMENT. THE TERMS OF THE AWARD NOTICE ARE INCORPORATED INTO THIS RESTRICTED STOCK UNIT AGREEMENT.

On the Grant Date specified in the Award Notice, Amgen Inc., a Delaware corporation (the “Company”), has granted to you, the grantee named in the Award Notice, an award under the Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan, as amended and/or restated from time to time (the “Plan”), for the Number of Units with respect to the number of shares of the \$0.0001 par value common stock of the Company (the “Shares”) specified in the Award Notice, on the terms and conditions set forth in this Restricted Stock Unit Agreement, any additional terms and conditions for your country set forth in the attached Appendix A and the Award Notice (collectively, the “Agreement”) and the Plan. The Units shall constitute Restricted Stock Units under Section 9.5 of the Plan, which is incorporated herein by reference. Capitalized terms not defined herein shall have the meanings assigned to such terms in the Plan.

I. Vesting Schedule and Termination of Units.

- a. *General.* Subject to the terms and conditions of this Agreement, on each Vesting Date, the Number of Units indicated on the Vesting Schedule shall vest, provided that you have remained continuously and actively employed with the Company or an Affiliate through each applicable Vesting Date, unless (i) [your employment has terminated due to your Voluntary Termination (as defined in paragraph (d) of this Section I below)]^{*2}, [(ii)] you experience a Qualified Termination (as defined below), or (iii) [(ii)] as otherwise determined by the Company in the exercise of its discretion as provided in paragraph (f) of this Section I. The Units represent an unfunded, unsecured promise by the Company to deliver Shares. Only whole Shares shall be issued upon vesting of the Units, and the Company shall be under no obligation to issue any fractional Shares to you. If your employment with the Company or an Affiliate is terminated for any reason or for no reason, including if your active employment is terminated by the Company or an Affiliate without Cause (as defined below), or in the event of any other termination of your active employment caused directly or indirectly by the Company or an Affiliate, except as otherwise provided in paragraphs (b), (c), [(d),]^{*2(1)} (e) or (f) of this Section I below, your unvested Units shall automatically expire and terminate on the date of termination of your active employment. Notwithstanding anything herein to the contrary, the Vesting Schedule

² Paragraph (d) of Section I of this Agreement is not applicable to awards identified by the Administrator as new hire, retention, special or promotion grants and the provisions of such paragraph shall be reserved and references thereto identified by an asterisk (*) shall be omitted from the agreements evidencing such grants.

may be accelerated (by notice in writing) by the Company in its sole discretion at any time that the Units remain outstanding and unvested (in whole or in part). In addition, if not prohibited by local law, vesting may be suspended by the Company in its sole discretion during a leave of absence as provided from time to time according to Company policies and practices.

- b. *Permanent and Total Disability.* Notwithstanding the provisions in paragraph (a) above, if your employment with the Company or an Affiliate terminates due to your Permanent and Total Disability (as defined below), then the vesting of Units granted under this Agreement shall be accelerated, subject to your execution and non-revocation of a general release and waiver in a form provided by the Company (for the purpose of resolving any potential or actual disputes arising from your employment and the termination of your employment with the Company) (a “Release”), to vest in full as of the day immediately preceding such termination of your employment with respect to all Units granted hereunder. Notwithstanding the foregoing, if the Units were granted in the calendar year in which such termination occurs, (i) the Units shall instead be accelerated to vest only with respect to a number of Units equal to (A) the number of Units subject to this Agreement multiplied by (B) a fraction, the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12), and (ii) any Units that remain unvested following the acceleration provided for in clause (i) shall automatically expire and terminate on the date of the termination of your active employment due to your Permanent and Total Disability without consideration therefor.
- c. *Death.* Notwithstanding the provisions in paragraph (a) above, if your employment with the Company or an Affiliate terminates due to your death, then the vesting of Units granted under this Agreement shall be accelerated to vest in full as of the day immediately preceding your death. Notwithstanding the foregoing, if the Units were granted in the calendar year in which your death occurs, (i) the Units shall instead be accelerated to vest only with respect to a number of Units equal to (A) the number of Units subject to this Agreement multiplied by (B) a fraction, the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your death occurs, and the denominator of which is twelve (12), and (ii) any Units that remain unvested following the acceleration provided for in clause (i) shall automatically expire and terminate on the date of the termination of your active employment due to your death without consideration therefor.
- d. *[Voluntary Termination (Retirement)].*
 - (i) Notwithstanding the provisions in paragraph (a) above, if you terminate your employment with the Company or an Affiliate due to your voluntary termination (and such voluntary termination is not the result of Permanent and Total Disability (as defined below)) after you are at least sixty-five (65) years of age, or after you are at least fifty-five (55) years of age and have been an employee of the Company and/or an Affiliate for at least ten (10) years in the aggregate as determined by the Company in

its sole discretion according to Company policies and practices as in effect from time to time (“Voluntary Termination”), then the Units will continue to vest following your Voluntary Termination pursuant to the Vesting Schedule without regard to the termination of employment prior to the Vesting Date, subject to your execution and non-revocation of a Release. Notwithstanding the foregoing, if the Units were granted in the calendar year in which the Voluntary Termination occurs, (i) the Units will continue to vest pursuant to the Vesting Schedule provided in the Award Notice, provided, that each tranche of Units scheduled to vest upon each remaining Vesting Date in the Vesting Schedule will vest only with respect to (A) the number of Units in such tranche multiplied by (B) a fraction, the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar in which your termination occurs and the denominator of which is twelve (12), and (ii) any Units in excess of the number of Units which are eligible to vest pursuant to clause (i) shall automatically expire and terminate on the date of termination of your active employment due to your Voluntary Termination without consideration therefor; provided, further, however, that in the event of your death following your Voluntary Termination, any Units that remain outstanding as of the date of your death will become vested (and the Vesting Date with respect to such Units will occur) as of the day immediately preceding your death.

(ii) Notwithstanding the foregoing, in the event your Voluntary Termination occurs on or after the date of a Change of Control, then, to the extent permitted by applicable law, the vesting of Units granted under this Agreement shall be accelerated to vest as of the day immediately prior to the date of your Voluntary Termination.

(iii) Notwithstanding the definition of Voluntary Termination set forth above, if the Company receives an opinion of counsel that there has been a legal judgment and/or legal development in your jurisdiction that would likely result in the favorable treatment upon Voluntary Termination described above being deemed unlawful and/or discriminatory, then the Committee will not apply the favorable treatment described above.][Reserved]*³

- e. *Qualified Termination after a Change of Control.* Notwithstanding the provisions in paragraph (a) above, in the event of your Qualified Termination (as defined below), then, to the extent permitted by applicable law, the vesting of Units granted under this Agreement shall be accelerated to vest as of the day immediately prior to the date of your Qualified Termination.
- f. *Continued Vesting.* Notwithstanding the provisions in paragraph (a) above, the Company may in its sole discretion at any time during the term of this Agreement, in writing, otherwise provide that the Units will vest pursuant to the Vesting Schedule without

³ Paragraph(d) of Section I of this Agreement is not applicable to awards identified by the Administrator as new hire, retention, special or promotion grants and the provisions of such paragraph shall be reserved and references thereto identified by an asterisk (*) shall be omitted from the agreements evidencing such grants.

regard to the termination of employment prior to the Vesting Date, subject to any terms and conditions that the Company may determine.

For purposes of this Agreement:

- (i) “termination of your active employment” shall mean the last date that you are either an active employee of the Company or an Affiliate or actively engaged as a Director of the Company or an Affiliate; in the event of termination of your employment (whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are working or the terms of your employment agreement, if any), your right to receive Units and vest under the Plan, if any, will terminate effective as of the date that you are no longer actively providing services and will not be extended by any notice period (*e.g.*, active employment would not include any period of “garden leave” or similar period mandated under employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any). The Company shall have exclusive discretion to determine when you are no longer actively providing services for purposes of this Agreement (including whether you may still be considered to be providing services while on a leave of absence);
- (ii) “Cause” shall mean (i) your conviction of a felony (or similar crime under applicable law, as determined by the Company), or (ii) your engaging in conduct that constitutes willful gross neglect or willful gross misconduct in carrying out your duties, resulting, in either case, in material economic harm to the Company or any Affiliate, unless you believed in good faith that such conduct was in, or not contrary to, the best interests of the Company or any Affiliate. For purposes of clause (ii) above, no act, or failure to act, on your part shall be deemed “willful” unless done, or omitted to be done, by you not in good faith;
- (iii) “Permanent and Total Disability” shall have the meaning ascribed to such term under Section 22(e)(3) of the Code and with such permanent and total disability being certified prior to termination of your employment by (i) the U.S. Social Security Administration, (ii) the comparable governmental authority applicable to an Affiliate, (iii) such other body having the relevant decision-making power applicable to an Affiliate, or (iv) an independent medical advisor appointed by the Company in its sole discretion, as applicable, in any such case;
- (iv) “Qualified Termination” shall mean
 - (a) if you are an employee who participates in the Change of Control Plan (as defined below), your termination of employment within two (2) years following a Change of Control (i) by the Company or an Affiliate other than for Cause, Disability, or as a result of your death or (ii) by you for Good Reason (as defined in the Change of Control Plan); or
 - (b) if you are an employee who does not participate in the Change of Control Plan or the Change of Control Plan is no longer in effect, your termination

of employment within two (2) years following a Change of Control by the Company or an Affiliate other than for Cause, Disability, or as a result of your death;

(v) “Change of Control” shall mean the occurrence of any of the following:

- (A) the acquisition (other than from the Company) by any person, entity or “group,” within the meaning of Section 13(d)(3) or 14(d)(2) of the Exchange Act (excluding, for this purpose, the Company or any of its Affiliates, or any employee benefit plan of the Company or any of its Affiliates which acquires beneficial ownership of voting securities of the Company), of beneficial ownership (within the meaning of Rule 13d-3 promulgated under the Exchange Act) of fifty percent (50%) or more of either the then-outstanding Shares or the combined voting power of the Company’s then-outstanding voting securities entitled to vote generally in the election of directors; or
- (B) the consummation by the Company of a reorganization, merger, consolidation, (in each case, with respect to which persons who were the stockholders of the Company immediately prior to such reorganization, merger or consolidation do not, immediately thereafter, own more than fifty percent (50%) of the combined voting power entitled to vote generally in the election of directors of the reorganized, merged or consolidated company’s then-outstanding voting securities) or a liquidation or dissolution of the Company or of the sale of all or substantially all of the assets of the Company.

Notwithstanding anything herein or in the Agreement to the contrary, if a Change of Control constitutes a payment event with respect to any Unit that is subject to United States income tax and which provides for a deferral of compensation that is subject to Section 409A (as defined below), the transaction or event described in subsection (A) or (B) above must also constitute a “change in control event,” as defined in U.S. Treasury Regulation § 1.409A-3(i)(5), in order to constitute a Change of Control for purposes of payment of such Unit.

- (vi) “Change of Control Plan” shall mean the Company’s change of control and severance plan, including the Amgen Inc. Change of Control Severance Plan, as amended and restated, effective as of December 9, 2010 (and any subsequent amendments thereto), or equivalent plan governing the provision of benefits to eligible employees upon the occurrence of a Change of Control (including resulting from a termination of employment that occurs within a specified time period following a Change of Control), as in effect immediately prior to a Change of Control; and
- (vii) “Disability” shall mean your “disability” as determined in accordance with the Company’s long-term disability plan as in effect immediately prior to a Change of Control.

II. Form and Timing of Settlement. Subject to satisfaction of Tax Obligations or similar obligations as provided for in Section III, any vested Units shall be settled by the Company delivering to you a number of Shares equal to the number of such vested Units or in a lump sum in cash with a value equal to the Fair Market Value of the number of Shares subject to the vested Units as of the applicable vesting date (without interest thereon), or in a combination of Shares and cash, as determined by the Administrator at any time prior to settlement and in its discretion. Such settlement of any vested Units shall occur on the earlier of (a) the regularly scheduled Vesting Date(s) indicated in the Vesting Schedule or in any case by the end of the calendar year in which such applicable Vesting Date occurs, or (b) the date of the applicable event set forth in paragraph (b), (c), [the last clause of paragraph (d)(i), (d)(ii),]* or (e) of Section I above or as soon as practicable, and in any event within 70 days after the date of such event. [(For the avoidance of doubt, in the event that any Units continue to vest following a Voluntary Termination in accordance with Section I(d)(i) above, the Vesting Date(s) for purposes of settlement pursuant to this Section II shall be the regularly scheduled Vesting Dates following such termination.)]*⁴ Notwithstanding the foregoing provisions of this Section II, to the extent that the Units are exempt from or otherwise not subject to Section 409A, the Units shall be settled within the short-term deferral period of Section 409A. Notwithstanding anything to the contrary in the foregoing, in the event that (i) the vesting and settlement of Units is conditioned on your execution, non-revocation and delivery of a Release and (ii) the period over which the Release becomes effective and nonrevocable commences in one calendar year and ends in the next calendar year, the Units will be settled in the second calendar year. Shares issued in respect of a Unit shall be deemed to be issued in consideration of past services actually rendered by you to the Company or an Affiliate or for its benefit for which you have not previously been compensated or for future services to be rendered, as the case may be, which the Company deems to have a value at least equal to the aggregate par value thereof.

III. Tax Withholding; Issuance of Shares. Regardless of any action the Company or your actual employer (the “Employer”) takes with respect to any or all income tax (including federal, state and local taxes), social insurance, payroll tax, fringe benefit tax, payment on account or other tax-related items related to your participation in the Plan and legally applicable to you (“Tax Obligations”), you acknowledge that the ultimate liability for all Tax Obligations is and remains your responsibility and may exceed the amount, if any, actually withheld by the Company and/or your Employer. You further acknowledge that the Company and/or your Employer (i) make no representations or undertakings regarding the treatment of any Tax Obligations in connection with any aspect of the Units or the underlying Shares, including the grant of the Units, the vesting of Units, the conversion of the Units into Shares or the receipt of an equivalent cash payment, the subsequent sale of any Shares acquired at vesting and the receipt of any Dividends (as defined in Section IV, below) or Dividend Equivalents, and (ii) do not commit to and are under no obligation to structure the terms of the grant or any aspect of the Units to reduce or eliminate your liability for Tax Obligations or achieve any particular tax result. Furthermore, if you become subject to tax in more than one jurisdiction, you acknowledge that the Company and/or your Employer (or former employer, as applicable) may be required to withhold or account for Tax Obligations in more than one jurisdiction.

⁴ Paragraph(d) of Section I of this Agreement is not applicable to awards identified by the Administrator as new hire, retention, special or promotion grants and the provisions of such paragraph shall be reserved and references thereto identified by an asterisk (*) shall be omitted from the agreements evidencing such grants.

Prior to any relevant taxable or tax withholding event, as applicable, you shall pay or make adequate arrangements satisfactory to the Company or to your Employer (in their sole discretion) to satisfy all Tax Obligations. In this regard, you authorize the Company and/or your Employer or their respective agents, at their discretion, to satisfy all applicable Tax Obligations by one or a combination of the following:

- (a) withholding from your wages or other cash compensation paid to you by the Company and/or your Employer; or
- (b) withholding from proceeds of the sale of Shares acquired upon vesting or payment of the Units either through your voluntary sale or through a mandatory sale arranged by the Company (on your behalf pursuant to this authorization); or
- (c) withholding in Shares issuable, or cash payable, upon vesting or payment of the Units, provided that, if such Shares are withheld, the Company and your Employer shall only withhold an amount of Shares with a fair market value not to exceed the Tax Obligations as determined in the discretion of the Company or your Employer, as applicable.

Depending on the withholding method, the Company may withhold or account for Tax Obligations by considering applicable minimum statutory withholding rates or other applicable withholding rates, including maximum applicable rates. If the Tax Obligations are satisfied by withholding in Shares, for tax purposes you are deemed to have been issued the full number of Shares subject to the vested Units, notwithstanding that a number of the Shares is held back and not actually issued to you solely for the purpose of paying the Tax Obligations due as a result of any aspect of your participation in the Plan (any Shares withheld by the Company hereunder shall not be deemed to have been issued by the Company for any purpose under the Plan and shall remain available for issuance thereunder).

Finally, you shall pay to the Company or your Employer any amount of Tax Obligations that the Company or your Employer may be required to withhold or account for as a result of your participation in the Plan that cannot be or were not satisfied by the means previously described. You agree to take any further actions and execute any additional documents as may be necessary to effectuate the provisions of this Section III. Notwithstanding Section II above, the Company may refuse to issue or deliver the Shares or the proceeds of the sale of Shares if you fail to comply with your obligations in connection with the Tax Obligations.

IV. Dividend Equivalents

- (a) Crediting and Payment of Dividend Equivalents. Subject to this Section IV, Dividend Equivalents shall be credited on each Unit granted to you under this Agreement in the manner set forth in the remainder of this Section IV. With respect to each Unit covered by the Award, if the Company declares one or more dividends or distributions (each, a “Dividend”) on its Common Stock with a record date which occurs during the period commencing on the Grant Date through and including the day immediately preceding the day the share of Common Stock subject to such Unit is issued to you, whether in the form

of cash, Common Stock or other property, then on the date such Dividend is paid to the Company's stockholders you shall be credited with an amount equal to the amount or fair market value of such Dividend which would have been payable to you if you held a share of Common Stock as of the record date for such Dividend, unless the applicable Unit has been forfeited between the record date and payment date for such Dividend. Any such Dividend Equivalents shall be credited and deemed reinvested in the Common Stock as of the applicable Dividend payment date. Dividend Equivalents shall be payable in full shares of Common Stock, unless the Administrator determines, at any time prior to payment and in its discretion, that they shall be payable in cash. Dividend Equivalents payable with respect to fractional shares of Common Stock shall be paid in cash.

- (b) Treatment of Dividend Equivalents. Except as otherwise expressly provided in this Section IV, any Dividend Equivalents credited to you shall be subject to all of the provisions of this Agreement which apply to the Unit with respect to which they have been credited and shall be payable, if at all, at the time and to the extent that the underlying Unit becomes payable pursuant to Section II above. Dividend Equivalents shall not be payable on any Units that do not vest, or are forfeited, pursuant to the terms of this Agreement. Dividend Equivalent rights and any amounts that may become distributable in respect thereof shall be treated separately from the Units and the rights arising in connection therewith for purposes of the designation of time and form of payments required by Section 409A of the Code (together with any Department of Treasury regulations and other interpretive guidance issued thereunder, including without limitation any such regulations or other guidance that may be issued after the Grant Date, "Section 409A").

V. Transferability. No benefit payable under, or interest in, this Agreement, the Units, or the Shares that are scheduled to be issued to you hereunder shall be subject in any manner to anticipation, alienation, sale, transfer, assignment, pledge, encumbrance or charge and any such attempted action shall be void and no such benefit or interest shall be, in any manner, liable for, or subject to, your or your beneficiary's debts, contracts, liabilities or torts; provided, however, nothing in this Section V shall prevent transfer (i) by will or (ii) by applicable laws of descent and distribution.

VI. Notices. Any notices provided for in this Agreement or the Plan shall be given in writing or electronically and shall be deemed effectively given upon receipt or, in the case of notices delivered by the Company to you, five (5) days after deposit in the United States mail or equivalent foreign postal service, postage prepaid, addressed to you at such address as is currently maintained in the Company's records or at such other address as you hereafter designate by written notice to the Company Stock Administrator. Such notices may be given using any automated system for the documentation, granting or settlement of Awards, such as a system using an internet website or interactive voice response, as approved by the Company.

VII. Plan. This Agreement is subject to all the provisions of the Plan, which provisions are hereby made a part of this Agreement, including without limitation the provisions of Section 9.5 of the Plan relating to Restricted Stock Units, and is further subject to all interpretations, amendments, rules and regulations which may from time to time be promulgated and adopted pursuant to the Plan. In the

event of any conflict between the provisions of this Agreement and those of the Plan, the provisions of the Plan shall control.

VIII. Governing Law and Venue. The terms of this Agreement shall be governed by the laws of the State of Delaware without giving effect to principles of conflicts of laws. For purposes of litigating any dispute that arises hereunder, the parties hereby submit to and consent to the jurisdiction of the State of Delaware, and agree that such litigation shall be conducted in the courts of the State of Delaware, or the federal courts for the United States for the federal district located in the State of Delaware, and no other courts, where this Agreement is made and/or to be performed.

IX. Code Section 409A. The time and form of payment of the Units is intended to comply with the requirements of Section 409A and this Agreement shall be interpreted in accordance with Section 409A. Accordingly, no acceleration or deferral of any payment shall be permitted if it would cause the payment of the Units to violate Section 409A. In addition, notwithstanding any provision herein to the contrary, in the event that following the Grant Date, the Committee (as defined in the Plan) determines that it may be necessary or appropriate to do so, the Committee may adopt such amendments to the Plan and/or this Agreement or adopt other policies and procedures (including amendments, policies and procedures with retroactive effect), or take any other actions, that the Committee determines are necessary or appropriate to (a) exempt the Plan and/or the Units from the application of Section 409A and/or preserve the intended tax treatment of the benefits provided with respect to this Award, or (b) comply with the requirements of Section 409A; provided, however, that this paragraph shall not create an obligation on the part of the Committee to adopt any such amendment, policy or procedure or take any such other action. For purposes of Section 409A, the right to receive payment of Units at each Vesting Date shall be treated as a right to receive separate and distinct payments. No payment hereunder shall be made to you during the six (6)-month period following your "separation from service" (within the meaning of Section 409A) to the extent that the Company determines that paying such amount at the time set forth herein would be a prohibited distribution under Section 409A(a)(2)(B)(i). If the payment of any such amounts is delayed as a result of the previous sentence, then within thirty (30) days following the end of such six (6)-month period (or, if earlier, your death), the Company shall pay to you (or to your estate) the cumulative amounts that would have otherwise been payable to you during such period, without interest.

X. Acknowledgement. By electing to accept this Agreement, you acknowledge receipt of this Agreement and hereby confirm your understanding that the terms set forth in this Agreement constitute, subject to the terms of the Plan, which terms shall control in the event of any conflict between the Plan and this Agreement, the entire agreement and understanding of the parties with respect to the matters contained herein and supersede any and all prior agreements, arrangements and understandings, both oral and written, between the parties concerning the subject matter of this Agreement. The Company may, in its sole discretion, decide to deliver any documents related to current or future participation in the Plan (including this Agreement) by electronic means. You hereby consent to receive such documents by electronic delivery and agree to participate in the Plan through an on-line or electronic system established and maintained by the Company or a third party designated by the Company.

XI. Acknowledgement of Nature of Plan and Units. In accepting this Agreement, you acknowledge, understand and agree that:

- (a) the Plan is established voluntarily by the Company, is discretionary in nature and may be modified, amended, suspended or terminated by the Company at any time, as provided in the Plan;
- (b) the Plan is operated and the Units are granted solely by the Company, and only the Company is a party to this Agreement; accordingly, any rights you may have under this Agreement may be raised only against the Company but not any Affiliate (including, but not limited to, the Employer);
- (c) no Affiliate (including, but not limited to, the Employer) has any obligation to make any payment of any kind to you under this Agreement;
- (d) the grant of the Units is exceptional, voluntary and occasional and does not create any contractual or other right to receive future awards of Units, or benefits in lieu of Units even if Units have been awarded in the past;
- (e) all decisions with respect to future awards, if any, will be at the sole discretion of the Company;
- (f) your participation in the Plan is voluntary;
- (g) the grant of Units, the Shares subject to the Units, and the income from and value of same, are not intended to replace any pension rights or compensation;
- (h) neither the grant of Units nor any provision of this Agreement, the Plan or the policies adopted pursuant to the Plan confer upon you any right with respect to employment or continuation of current employment and shall not interfere with the ability of your Employer to terminate your employment or service relationship (if any) at any time;
- (i) in the event that you are not an employee of the Company or any Affiliate, the Units shall not be interpreted to form an employment contract or relationship with the Company or any Affiliate;
- (j) the future value of the underlying Shares is unknown, indeterminable and cannot be predicted with certainty;
- (k) in consideration of the grant of Units hereunder, no claim or entitlement to compensation or damages arises from termination of Units, and no claim or entitlement to compensation or damages shall arise from forfeiture of the Units resulting from (i) the termination of your employment by the Company or an Affiliate (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any)

and/or (ii) the application of the Clawback Policy as described in Section XVIII below, and you irrevocably release the Company and your Employer from any such claim that may arise; if, notwithstanding the foregoing, any such claim is found by a court of competent jurisdiction to have arisen, you shall be deemed irrevocably to have waived your entitlement to pursue such claim;

(l) unless otherwise agreed with the Company, the Units, the Shares subject to the Units, and the income from and value of same, are not granted as consideration for, or in connection with, the service you may provide as a director of an Affiliate of the Company;

(m) except as otherwise provided in this Agreement or the Plan, the Units and the benefits evidenced by this Agreement do not create any entitlement to have the Units or any such benefits transferred to, or assumed by, another company nor to be exchanged, cashed out or substituted for, in connection with any corporate transaction affecting the shares of the Company;

(n) the following provisions apply only if you are providing services outside the United States:

(i) for employment law purposes outside the United States, the Units, Shares subject to the Units, and the income from and value of same, are not part of normal or expected compensation or salary for any purpose, including but not limited to for purposes of calculating any severance, resignation, termination, redundancy, dismissal, end of service payments, bonuses, holiday pay, long-service awards, pension or retirement benefits or similar mandatory payments; and

(ii) neither the Company, your Employer nor any Affiliate of the Company shall be liable for any foreign exchange rate fluctuation between your local currency and the United States Dollar that may affect the value of the Units or of any amounts due to you pursuant to the settlement of the Units or the subsequent sale of any Shares acquired upon settlement.

XII. No Advice Regarding Award. The Company is not providing any tax, legal or financial advice, nor is the Company making any recommendations regarding your participation in the Plan, or your acquisition or sale of the underlying Shares. You should consult with your own personal tax, legal and financial advisors regarding your participation in the Plan before taking any action related to the Plan.

XIII. Compliance with Laws. Notwithstanding any provision of this Agreement to the contrary, if you are employed by the Company or an Affiliate in any of the countries identified in the attached Appendix A (which constitutes a part of this Agreement), are subject to the laws of any foreign jurisdiction, or relocate to one of the countries included in the attached Appendix A, the Units granted hereunder shall be subject to any additional terms and conditions for your country set forth in Appendix A and to the following additional terms and conditions:

(a) the terms and conditions of this Agreement, including Appendix A, are deemed modified to the extent necessary or advisable to comply with applicable foreign laws or facilitate the administration of the Plan;

- (b) if applicable, the effectiveness of your award of Units is conditioned upon its compliance with any applicable foreign laws, regulations, rules or local governmental regulatory exemption and subject to receipt of any required foreign regulatory approvals;
- (c) to the extent necessary to comply with applicable foreign laws, the payment of any earned Units shall be made in cash or Common Stock, at the Company's election; and
- (d) the Company may take any other action, before or after an award of Units is made, that it deems advisable to obtain approval or comply with any necessary local governmental regulatory exemptions or approvals.

Notwithstanding anything to the contrary contained herein, the Company shall not take any actions hereunder that would violate the Securities Act, the Exchange Act, the Code, or any other securities or tax or other applicable law or regulation, or the rules of any Securities Exchange. Notwithstanding anything to the contrary contained herein, the Shares issuable upon vesting of the Unit shall not be issued unless such Shares are then registered under the Securities Act, or, if such Shares are not then so registered, the Company has determined that such vesting and issuance would be exempt from the registration requirements of the Securities Act, and that the issuance satisfied all other applicable legal requirements.

XIV. Data Privacy. *In order for the Company to facilitate your participation in the Plan, the Company and your Employer must collect and use personal data about you. In accordance with applicable laws, reasonable security measures will be implemented and maintained to protect the security of your personal data; however, you understand that absolute security cannot be guaranteed.*

You understand that the Company and your Employer may hold certain personal information about you, including your name, home address and telephone number, email address, date of birth, social insurance/security number (to the extent permitted under applicable local law), passport or other identification number, salary, nationality, job title/work history/service periods, residency status, citizenship, tax withholding and payroll data, any shares of stock or directorships held in the Company, details of all equity compensation or any other entitlement to Shares awarded, cancelled, vested, unvested or outstanding in your favor, for the purposes of implementing, administering and managing the Plan ("personal data").

You authorize the transfer of your personal data to Merrill Lynch Bank & Trust Co., FSB, or any successor thereto, and any other third parties which may assist the Company (presently or in the future) with implementing, administering and managing your participation in the Plan to receive, possess, use, retain and transfer your personal data, in electronic or other form, for the purpose of implementing, administering and managing your participation in the Plan, including any requisite transfer of such personal data as may be required to any other broker, escrow agent or other third party with whom the Shares received in settlement of the Units may be deposited. You understand that such authorized recipients of your personal data may be located in countries that do not provide the same level of data privacy laws and protections as the country in which your personal data originated. Transfers of personal data among Company and its group entities follow applicable laws and our Binding Corporate Rules (BCRs). For more information on Company's BCRs, please visit <http://www.amgen.com/bcr/>. You acknowledge that the collection, use and transfer of your personal

data is necessary to facilitate your participation in the Plan, as well as to grant you Units or other equity awards and administer or maintain such awards.

You may correct or update your personal data previously provided to Company, by contacting your local human resources representative. Subject to applicable law, you may have additional rights, including the right to object and/or request destruction of your personal data. To exercise these rights, where applicable, please contact your local human resources representative.

XV. Severability. If one or more of the provisions of this Agreement shall be held invalid, illegal or unenforceable in any respect, the validity, legality and enforceability of the remaining provisions shall not in any way be affected or impaired thereby and the invalid, illegal or unenforceable provisions shall be deemed null and void; however, to the extent permissible by law, any provisions which could be deemed null and void shall first be construed, interpreted or revised retroactively to permit this Agreement to be construed so as to foster the intent of this Agreement and the Plan.

XVI. Language. By electing to accept this Agreement, you acknowledge that you are sufficiently proficient in English, or have consulted with an advisor who is sufficiently proficient in English, so as to allow you to understand the terms and conditions of this Agreement. Further, if you have received this Agreement or any other document related to the Plan translated into a language other than English and if the meaning of the translated version is different than the English version, the English version will control, unless otherwise required by applicable law.

XVII. Imposition of Other Requirements. The Company reserves the right to impose other requirements on your participation in the Plan, on the Units and on any Shares acquired under the Plan, to the extent the Company determines it is necessary or advisable for legal or administrative reasons, and to require you to sign any additional agreements or undertakings that may be necessary to accomplish the foregoing.

XVIII. Compensation Subject to Recovery. The Units subject to and Shares issuable under this Award and all compensation payable with respect to them shall be subject to clawback, recoupment and/or recovery by the Company pursuant to any and all of the Company's policies with respect to the clawback, recoupment or recovery of compensation in effect as of the Grant Date or as may be adopted or maintained by the Company following the Grant Date, including, without limitation, the Company's Policy on Recovery of Erroneously Awarded Compensation (effective October 2, 2023) and Executive Officer Equity Recoupment Policy (effective December 31, 2020), as they shall be in effect and may be amended from time to time (collectively, the "Clawback Policy"), to the maximum extent permitted by applicable law.

XIX. Waiver. You acknowledge that a waiver by the Company of breach of any provision of this Agreement shall not operate or be construed as a waiver of any other provision of this Agreement, or of any subsequent breach by you or any other grantee.

XX. Headings. This Agreement's section headings are for convenience only and shall not constitute a part of this Agreement or affect this Agreement's meaning.

Very truly yours,
AMGEN INC.

By: _____
Name:
Title:

APPENDIX A

ADDITIONAL TERMS AND CONDITIONS OF THE AMGEN INC. SECOND AMENDED AND RESTATED 2009 EQUITY INCENTIVE PLAN, AS AMENDED AND/OR RESTATED FROM TIME TO TIME

GRANT OF RESTRICTED STOCK UNITS (BY COUNTRY)

Certain capitalized terms used but not defined in this Appendix A shall have the meanings set forth in the Plan and/or the Agreement to which this Appendix is attached.

TERMS AND CONDITIONS

This Appendix includes additional terms and conditions that govern any Units granted under the Plan if, under applicable law, you are a resident of, are deemed to be a resident of or are working in one of the countries listed below. Furthermore, the additional terms and conditions that govern any Units granted hereunder may apply to you if you transfer employment and/or residency to one of the countries listed below and the Company shall, in its discretion, determine to what extent the terms and conditions contained herein shall apply to you.

NOTIFICATIONS

This Appendix also includes notifications relating to exchange control and other issues of which you should be aware with respect to your participation in the Plan. The information is based on the exchange control, securities and other laws in effect in the countries to which this Appendix refers as of October 2025. Such laws are often complex and change frequently. As a result, the Company strongly recommends that you not rely on the notifications herein as the only source of information relating to the consequences of your participation in the Plan because the information may be outdated when you vest in the Units and acquire Shares under the Plan, or when you subsequently sell Shares acquired under the Plan.

In addition, the notifications are general in nature and may not apply to your particular situation, and the Company is not in a position to assure you of any particular result. Accordingly, you should seek appropriate professional advice as to how the relevant laws in your country may apply to your situation. Finally, if you are a citizen or resident of a country other than the one in which you are currently residing and/or working or are considered a resident of another country for local law purposes, the information contained herein may not be applicable to you or you may be subject to the provisions of one or more jurisdictions.

ALL NON-U.S. JURISDICTIONS

TERMS AND CONDITIONS

Tax Withholding; Issuance of Shares. The following provision supplements Section III of the Agreement:

In the event the Company withholds or accounts for Tax Obligations by considering maximum applicable rates in your jurisdiction(s), in the event of over-withholding, you may receive a refund of any over-withheld amount in cash and will not be entitled to the equivalent amount in Shares, or if not refunded, you may seek a refund from the local tax authorities. In the event of under-withholding, you may be required to pay any additional Tax Obligations directly to the applicable tax authority or to the Company and/or your Employer.

NOTIFICATIONS

Insider Trading Restrictions/Market Abuse Laws. You may be subject to insider trading restrictions and/or market abuse laws based on the exchange on which the Shares are listed and in applicable jurisdictions including the United States and your country or your broker's country, if different, which may affect your ability to accept, acquire, sell or otherwise dispose of Shares, rights to Shares (e.g., Units) or rights linked to the value of Shares (e.g., Dividend Equivalents) during such times as you are considered to have "inside information" regarding the Company (as defined by the laws in applicable jurisdictions). Local insider trading laws and regulations may prohibit the cancellation or amendment of orders you place before you possessed inside information. Furthermore, you could be prohibited from (i) disclosing the inside information to any third party, which may include fellow employees (other than on a "need to know" basis) and (ii) "tipping" third parties or causing them otherwise to buy or sell securities. Any restrictions under these laws or regulations are separate from and in addition to any restrictions that may be imposed under any applicable Company insider trading policy. You are responsible for ensuring your compliance with any applicable restrictions and you should speak with your personal legal advisor on this matter.

Foreign Asset/Account, Tax Reporting Information. Your country of residence may have certain foreign asset and/or account reporting requirements which may affect your ability to acquire or hold Shares under the Plan or cash received from participating in the Plan (including from any Dividends or Dividend Equivalents received, or sale proceeds arising from the sale of Shares) in a brokerage or bank account outside of your country. You may be required to report such accounts, assets or transactions to the tax or other authorities in your country. You also may be required to repatriate sale proceeds or other funds received as a result of participating in the Plan to your country within a certain time after receipt. You are responsible for ensuring your compliance with such regulations, and you should speak with your personal legal advisor on this matter.

ALL EUROPEAN ECONOMIC AREA (“EEA”) / EUROPEAN UNION (“EU”) JURISDICTIONS, UNITED KINGDOM AND SWITZERLAND

TERMS AND CONDITIONS

Data Privacy Notice. This provision replaces Section XIV of the Agreement:

Please refer to the Fair Processing Notice previously provided by your local human resources representative, which notice governs the collection, use and transfer of your personal data necessary for the Company to facilitate your participation in the Plan. If you have any questions or concerns regarding the Fair Processing Notice, including questions about your rights afforded thereunder, you should contact your local human resources representative or send an email to hrconnect@amgen.com.

For purposes of implementing, administering and managing the Plan, Company and your Employer may hold certain personal data about you, including your name, home address and telephone number, email address, date of birth, social insurance/security number (to the extent permitted under applicable local law), passport or other identification number, salary, nationality, job title/work history/service periods, residency status, citizenship, tax withholding and payroll data, any shares of stock or directorships held in the Company, details of all equity compensation or any other entitlement to Shares awarded, cancelled, vested, unvested or outstanding in your favor (“personal data”).

You authorize the transfer of your personal data to Merrill Lynch Bank & Trust Co., FSB, or any successor thereto, and any other third parties which may assist the Company (presently or in the future) with implementing, administering and managing your participation in the Plan to receive, possess, use, retain and transfer your personal data, in electronic or other form, for the purpose of implementing, administering and managing your participation in the Plan, including any requisite transfer of such personal data as may be required to any other broker, escrow agent or other third party with whom the Shares received in settlement of the Units may be deposited.

ARGENTINA

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section XI of the Agreement:

In accepting this Agreement, you acknowledge, understand and agree that the grant of the Units is made by the Company (not your Employer) in its sole discretion and that the value of the Units or any Shares acquired under the Plan shall not constitute salary or wages for any purpose under Argentine labor law including, but not limited to, the calculation of (i) any labor benefits including, without limitation, vacation pay, thirteenth salary, compensation in lieu of notice, annual bonus, disability, and leave of absence payments, etc., or (ii) any termination or severance indemnities or similar payments.

NOTIFICATIONS

Securities Law Information. The Units and the underlying Shares have not been and will not be publicly issued, placed, distributed, offered or listed in the Argentine capital markets, and, as a result, have not been and will not be registered with the Argentina Securities Commission (Comisión Nacional de Valores, “CNV”). Neither this nor any other offering material related to the Units nor the underlying Shares may be utilized in connection with any general offering to the public in Argentina. Any Argentine resident who acquires Shares under the Plan does so under their own responsibility under the terms of a private offering to the Argentine resident from outside Argentina. Any Argentine resident who acquires Shares shall not transfer such Shares to any other person within six (6) months of acquiring the Shares, unless the transaction is conducted outside Argentina.

Exchange Control Information. Exchange control regulations in Argentina are subject to frequent change. You should consult with your personal legal advisor regarding any exchange control obligations that you may have prior to receiving proceeds from Dividend Equivalents, the sale of Shares or dividends. You must comply with any and all Argentine currency exchange restrictions, approvals and reporting requirements in connection with your participation in the Plan.

Foreign Asset/Account Reporting Information. If you are an Argentine resident, you are required to report certain information regarding any Shares you hold as of December 31 each year to the Argentine tax authorities on your annual tax return.

AUSTRALIA

NOTIFICATIONS

Australia Offer Document. This grant of Units is being made under Division 1A, Part 7.12 of the Corporations Act 2001 (Cth).

Tax Information. Subdivision 83A-C of the Income Tax Assessment Act 1997 (Cth) applies to the Units granted under the Plan, such that the Units are intended to be subject to deferred taxation.

Exchange Control Information. If you are an Australian resident, exchange control reporting is required for cash transactions exceeding a certain threshold (currently AUD10,000) and for international fund transfers. If an Australian bank is assisting with the transaction, the bank will file the report on your behalf. If there is no Australian bank involved in the transfer, you will be required to file the report.

AUSTRIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If you are an Austrian resident and you hold Shares acquired under the Plan outside of Austria, you may be subject to reporting obligations to the Austrian National Bank.

Exchange Control Information. If the value of Shares meets or exceeds a certain threshold (currently, €5,000,000), you must report the securities held on a quarterly basis to the Austrian National Bank as of the last day of the quarter, on or before the 15th day of the month following the end of the calendar quarter. Further, if you hold cash in accounts outside Austria, monthly reporting requirements will apply, as explained in the next paragraph, if the aggregate transaction volume of such cash accounts meets or exceeds a certain threshold.

A separate reporting requirement applies when you sell Shares acquired under the Plan, receive a cash Dividend paid on such Shares or Dividend Equivalents paid in cash. In that case, there may be exchange control obligations if the cash proceeds are held outside of Austria. If the transaction volume of all cash accounts abroad meets or exceeds a certain threshold (currently €10,000,000), the movements and balances of all accounts must be reported monthly, as of the last day of the month, on or before the 15th day of the following month, on the prescribed forms.

BELGIUM

NOTIFICATIONS

Tax Reporting; Foreign Asset/Account Reporting Information. If you are a Belgian resident, you are required to report any securities (e.g., Shares acquired under the Plan) held and bank accounts (including brokerage accounts) opened and maintained outside of Belgium on your annual tax return. The first time you report the foreign security and/or bank account on your annual income tax return you will have to provide the National Bank of Belgium Central Contact Point with the account details of any such foreign accounts (including the account number, bank name and country in which such account was opened) in a separate form. This report, as well as information on how to complete it, can be found on the website of the National Bank of Belgium, www.nbb.be, under the *Kredietcentrales / Centrales des crédits* caption.

Stock Exchange Tax Information. A stock exchange tax applies to transactions executed by a Belgian resident through a non-Belgian financial intermediary, such as a U.S. broker. The stock exchange tax likely will apply when Shares acquired under the Plan are sold. It is your responsibility to comply with this tax obligation and you should consult your personal tax advisor for additional details on your obligations with respect to the stock exchange tax.

Annual Securities Accounts Tax Information. An annual securities accounts tax may be payable if the total value of securities held in a Belgian or foreign securities account (e.g., Shares acquired under the Plan) exceeds a certain threshold on four reference dates within the relevant reporting period (i.e., December 31, March 31, June 30 and September 30). In such case, the tax will be due on the value of the qualifying securities held in such account. It is your responsibility to comply with this obligation, and you should consult with your personal tax or financial advisor for additional details.

BRAZIL

TERMS AND CONDITIONS

Compliance with Law. By accepting the Units, you acknowledge that you agree to comply with applicable Brazilian laws and pay any and all applicable taxes associated with the vesting of the Units, the sale of Shares acquired under the Plan, the payment of Dividends on such Shares and the receipt of any Dividend Equivalents paid in cash.

Acknowledgement of Nature of Plan and Units. This provision supplements Section XI of the Agreement:

In accepting this Agreement, you acknowledge (i) that you are making an investment decision, (ii) that the Shares will be issued to you only if the vesting conditions are met and any necessary services are rendered by you during the vesting period set forth in the Vesting Schedule, and (iii) that the value of the underlying Shares is not fixed and may increase or decrease in value over the vesting period without compensation to you.

NOTIFICATIONS

Exchange Control Information. If you are resident or domiciled in Brazil, you will be required to submit annually a declaration of assets and rights held outside of Brazil to the Central Bank of Brazil if the aggregate value of such assets and rights on December 31 of each year exceeds US\$1,000,000. If such amount exceeds US\$100,000,000, the referenced declaration must be submitted quarterly, in the month following the end of each quarter. Assets and rights that must be reported include the following: (i) bank deposits; (ii) loans; (iii) financing transactions; (iv) leases; (v) direct investments; (vi) portfolio investments, including Shares acquired under the Plan; (vii) financial derivatives investments; and (viii) other investments, such as real estate. Please note that foreign individuals holding Brazilian visas are considered Brazilian residents for purposes of this reporting requirement and must declare at least the assets held abroad that were acquired subsequent to the date of admittance as a resident of Brazil. Individuals holding assets and rights outside of Brazil valued at less than US\$1,000,000 are not required to submit a declaration.

Tax on Financial Transactions. Payments to foreign countries, repatriation of funds into Brazil, and the conversion associated with such fund transfers into local currency, may be subject to the Tax on Financial Transactions. It is your responsibility to comply with any applicable Tax on Financial Transactions arising from participation in the Plan and you should consult your personal tax advisor for additional details.

BULGARIA

Exchange Control Information. You will be required to file statistical forms with the Bulgarian National Bank annually regarding your receivables in bank accounts abroad as well as securities held abroad (e.g., Shares acquired under the Plan) if the total sum of all such receivables and securities equals or exceeds a certain threshold as of the previous calendar year-end. The reports are due by

March 31. You should contact your bank in Bulgaria for additional information regarding these requirements.

CANADA

TERMS AND CONDITIONS

Termination of Employment. Section I(i) of the Agreement is amended to read as follows:

- (i) “termination of your active employment” shall mean the last date that you are either actually providing services to the Company or an Affiliate or actually providing services as a Director of the Company or an Affiliate; in the event of involuntary termination of your employment (regardless of the reason for such termination and whether or not later found to be invalid or unlawful, including for breaching applicable laws in the jurisdiction where you are employed or the terms of your employment agreement, if any), except as explicitly and minimally required under applicable legislation your right, if any, to earn, seek damages in lieu of, vest in or otherwise benefit from or participate in any portion of the Units or the Plan will be measured by and immediately terminate, as of the termination of your employment (the “Termination Date”). Except as explicitly and minimally required under applicable legislation, the Termination Date will exclude and will not be extended by any period during which notice, pay in lieu of notice or related payments or damages are provided or required to be provided under applicable laws (including, but not limited to, statute, contract, common/civil law or otherwise). You will not earn or be entitled to any pro-rated vesting of the Units or other benefits under or participation in the Plan for that portion of time before the Termination Date, nor will you be entitled to any compensation for lost vesting, benefits, or other participation. Subject to applicable legislation, if the date you are no longer actually providing services cannot be reasonably determined under the terms of this Agreement or the Plan, the Company shall have the exclusive discretion to determine when you are no longer actually providing services for purposes of this Agreement (including whether you may still be considered to be providing services while on a leave of absence). Notwithstanding the foregoing, if applicable employment standards legislation explicitly requires continued vesting or other participation during a statutory notice period, your right to vest in the Units or otherwise participate in or benefit from the Units under the Plan, if any, will terminate effective as of the last day of your minimum statutory notice period, but you will not earn or be entitled to pro-rated vesting or other benefits or participation if the vesting date falls after the end of your statutory notice period, nor will you be entitled to any compensation for lost vesting benefits, or participation;

Acknowledgement of Nature of Plan and Units. Sections XI(c), XI(i) and XI(k) of the Agreement are amended to read as follows:

- (c) no Affiliate (including, but not limited to, the Employer) has any obligation to make any payment of any kind to you under this Agreement, except to the extent explicitly and minimally required under applicable legislation;

(i) in consideration of the grant of Units hereunder, no claim or entitlement to compensation or damages arises from termination of Units, and no claim or entitlement to compensation or damages shall arise from forfeiture of the Units resulting from (i) the termination of your employment by the Company or an Affiliate (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any) and/or (ii) the application of the Clawback Policy as described in Section XVIII below, except to the extent explicitly and minimally required under applicable legislation;

(k) for employment law purposes outside the United States, the Units, Shares subject to the Units, and the income from and value of same, are not part of normal or expected compensation or salary for any purpose, including but not limited to for purposes of calculating any severance, resignation, termination, redundancy, dismissal, end of service payments, bonuses, holiday pay, long-service awards, pension or retirement benefits or similar mandatory payments, except to the extent explicitly and minimally required under applicable legislation;

Form of Settlement – Units Payable Only in Shares. Notwithstanding any discretion in Section 9.5 of the Plan or anything to the contrary in the Agreement, the Units do not provide any right for you, as a resident of Canada, to receive a cash payment and shall be paid in Shares only.

The following provision will apply to you if you are a resident of Quebec:

French Language Documents. A French translation of this document and certain other documents related to this Award will be made available to you as soon as reasonably practicable. You understand that, from time to time, additional information related to the Award may be provided in English and such information may not be immediately available in French. However, upon request, the Company will provide a translation of such information into French as soon as reasonably practicable. Notwithstanding anything to the contrary in the Agreement, and unless you indicate otherwise, the French translation of this document and certain other documents related to the Award will govern your participation in the Plan.

Data Privacy Notice. This provision supplements Section XIV of the Agreement:

You hereby authorize the Company and the Company's representative to discuss with and obtain all relevant information from all personnel (professional or not) involved in the administration of the Plan. You further authorize the Company, your Employer and Merrill Lynch Bank & Trust Co., FSB (or any other stock plan service provider) to disclose and discuss your participation in the Plan with their advisors. You also authorize the Company and your Employer to record such information and keep it in your file.

NOTIFICATIONS

Securities Law Information. You are permitted to sell Shares acquired through the Plan through the designated broker appointed under the Plan, if any, provided that the resale of such Shares takes place outside of Canada through the facilities of a stock exchange on which the Shares are listed (*e.g.*, the Nasdaq Global Select Market).

Foreign Asset/Account Reporting Information. Specified foreign property, including Shares, stock options and other rights to receive Shares (*e.g.*, Units) of a non-Canadian company held by a Canadian resident employee generally must be reported annually on a Form T1135 (Foreign Income Verification Statement) if the total cost of the employee's specified foreign property exceeds a certain threshold (currently C\$100,000) at any time during the year. Thus, such stock options and Units must be reported – generally at nil cost – if the applicable cost threshold is exceeded because other specified foreign property is held by the employee. When Shares are acquired, their cost generally is the adjusted cost base ("ACB") of the Shares. The ACB ordinarily would equal the fair market value of the Shares at the time of acquisition, but if the employee owns other shares of the same company, this ACB may have to be averaged with the ACB of the other shares.

CHINA

TERMS AND CONDITIONS

The following terms apply only to individuals who are subject to exchange control restrictions in the People's Republic of China (the "PRC"), as determined by the Company in its sole discretion:

Vesting of the Units. [Notwithstanding anything to the contrary in Section I(d) of the Agreement, if your employment with the Company or an Affiliate terminates due to your Voluntary Termination, as defined in Section I(d), then the vesting of Units granted under this Agreement shall be accelerated to vest as of the day immediately preceding such Voluntary Termination with respect to all Units granted hereunder.]*⁵

Sale Requirement. Notwithstanding anything to the contrary in the Agreement, due to exchange control laws in the PRC, you agree that the Company reserves the right to require the immediate sale of any Shares issued upon settlement of the Units. You understand and agree that any such immediate sale of Shares will occur as soon as is practical following settlement of the Units. Alternatively, if the Shares are not immediately sold upon settlement of the Units, the Company will require the sale of any Shares you may then hold within six (6) months (or such other period as may be required under applicable legal or exchange control requirements) following the termination of your employment with the Company, including its Affiliates.

You agree that the Company is authorized to instruct Merrill Lynch Bank & Trust Co., FSB or such other designated broker as may be selected by the Company to assist with the sale of the

⁵ Paragraph (d) of Section I of the Agreement is not applicable to awards identified by the Administrator as new hire, retention, special or promotion grants and the provisions of such paragraph shall be reserved and references thereto identified by an asterisk (*) shall be omitted from the agreements evidencing such grants.

Shares on your behalf pursuant to this authorization, and you expressly authorize such broker to complete the sale of such Shares. You also agree to sign any agreements, forms and/or consents that may be reasonably requested by the Company (or the Company's designated broker) to effectuate the sale of the Shares (including, without limitation, as to the transfers of the proceeds and other exchange control matters noted below) and to otherwise cooperate with the Company with respect to such matters, provided that you shall not be permitted to exercise any influence over how, when or whether the sales occur. Upon the sale of the Shares, you will receive the cash proceeds from the sale, less any applicable Tax Obligations, brokerage fees or commissions, in accordance with applicable exchange control laws and regulations.

You acknowledge that Merrill Lynch Bank & Trust Co., FSB or such other designated broker as may be selected by the Company is under no obligation to arrange for the sale of the Shares at any particular price. Due to fluctuations in the Share price and/or applicable exchange rates between the settlement date and (if later) the date on which the Shares are sold, the amount of proceeds ultimately distributed to you may be more or less than the market value of the Shares on the settlement date (which is the amount relevant to determining your liability for Tax Obligations). You understand and agree that the Company is not responsible for the amount of any loss that you may incur and that the Company assumes no liability for any fluctuations in the Share price and/or any applicable exchange rate.

Designated Broker Account. If Shares issued upon the settlement of the Units are not immediately sold, you acknowledge that you are required to maintain the Shares in an account with Merrill Lynch Bank & Trust Co., FSB or such other designated broker as may be selected by the Company until the Shares are sold through such Company-designated broker.

Exchange Control Requirements. You understand and agree that, pursuant to local exchange control requirements, you will be required to repatriate the cash proceeds from the sale of the Shares issued to you upon settlement of the Units and from the receipt of any Dividends or Dividend Equivalents to China. You further understand that, under applicable laws, such repatriation of your cash proceeds will need to be effectuated through a special exchange control account established by the Company or any Affiliate, including your Employer, and you hereby consent and agree that any proceeds may be transferred to such special account prior to being delivered to you. You also understand that the Company will deliver the proceeds to you as soon as possible, but that there may be delays in distributing the funds to you due to exchange control requirements in China. Proceeds may be paid to you in U.S. dollars or local currency at the Company's discretion. If the proceeds are paid to you in U.S. dollars, you will be required to set up a U.S. dollar bank account in China so that the proceeds may be deposited into this account. If the proceeds are paid to you in local currency, the Company is under no obligation to secure any particular currency conversion rate and the Company may face delays in converting the proceeds to local currency due to exchange control restrictions. You further agree to comply with any other requirements that may be imposed by the Company in the future in order to facilitate compliance with exchange control requirements in China.

COLOMBIA

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section XI of the Agreement:

You acknowledge that pursuant to Article 15 of Law 50/1990 (Article 128 of the Colombian Labor Code), the Plan and related benefits do not constitute a component of “salary” for any purpose. Therefore, they are considered to be of an extraordinary nature and will not be included and/or considered for purposes of calculating any and all labor benefits, such as legal/fringe benefits, vacations, indemnities, payroll taxes, social insurance contributions and/or any other labor-related amounts, subject to the limitations provided in Law 1393/2010.

Mandate Letter. In accepting the Units, you agree that – if requested by the Company or the Employer – you will execute a Mandate Letter or such other document (whether electronically or by such other method as requested by the Company or the Employer) that the Company determines is necessary or advisable in order that (i) a sufficient number of Shares to be allocated to you upon vesting can be sold on your behalf to cover Tax Obligations required to be withheld by the Employer and (ii) the proceeds from such sale can be wired directly from the Company to the Employer in Colombia for remittance to the tax authorities.

NOTIFICATIONS

Securities Law Information. The Shares are not and will not be registered with the Colombian registry of publicly traded securities (*Registro Nacional de Valores y Emisores*) and therefore the Shares may not be offered to the public in Colombia. Nothing in this document should be construed as the making of a public offer of securities in Colombia.

Exchange Control Information. Investment in assets located abroad (such as Shares acquired under the Plan) does not require prior approval from the Central Bank (*Banco de la República*). Nonetheless, such investments are subject to registration before the Central Bank as foreign investments held abroad, regardless of value.

Any payments for your investment originating in Colombia (and the liquidation of such investments) must be transferred through the Colombian foreign exchange market (*e.g.*, local banks), which includes the obligation to correctly complete and file the appropriate foreign exchange form (*declaración de cambio*).

Foreign Asset/Account Reporting Notice. An annual information return may need to be filed with the Colombian Tax Office detailing any assets held abroad (including Shares acquired under the Plan). If the individual value of any of these assets exceeds a certain threshold, each asset must be described (*e.g.*, its nature and its value) and the jurisdiction in which it is located must be disclosed. It is your responsibility to comply with this tax reporting requirement.

CROATIA

NOTIFICATIONS

Exchange Control Information. Croatian residents may be required to report any foreign investments (including Shares acquired under the Plan) to the Croatian National Bank for statistical purposes. You should be aware that exchange control regulations in Croatia are subject to frequent change and you are solely responsible for ensuring your continued compliance with current Croatian exchange control laws.

CZECH REPUBLIC

NOTIFICATIONS

Exchange Control Information. If you are a resident of the Czech Republic, you may be required to notify the Czech National Bank (“**CNB**”) of the acquisition of Shares under the Plan or maintenance of a foreign account if (i) you maintain foreign direct investments with a value of 2,500,000 Kč or more in the aggregate, (ii) you maintain a certain threshold of foreign financial assets, or (iii) you are specifically requested to do so by the CNB.

DENMARK

TERMS AND CONDITIONS

Danish Stock Option Act. In accepting the Units, you acknowledge that you have received an Employer Statement translated into Danish, which is being provided to comply with the Danish Stock Option Act. To the extent more favorable to you and required to comply with the Stock Option Act, as amended with effect from January 1, 2019. You also acknowledge any grant of Units under the Plan made on or after January 1, 2019 is subject to the rules of the amended Act. Accordingly, you agree that the treatment of the Units upon the termination of your employment is governed solely by Section I(i) of the Agreement and any corresponding provisions in the Plan. The relevant termination provisions are also detailed in the Employer Statement.

Please be aware that as set forth in Section 1 of the Act, the Act only applies to “employees” as that term is defined in Section 2 of the Act. If you are a member of the registered management of an Affiliate or Subsidiary in Denmark or otherwise do not satisfy the definition of employee, you will not be subject to the Act, and the Employer Statement will not apply to you.

NOTIFICATIONS

Foreign Asset/Account Reporting Information. The requirement to report certain information to the Danish Tax Administration via Form V or K was eliminated effective January 1, 2019. However, you still must report the foreign bank/brokerage accounts and their deposits, and Shares held in a foreign bank or brokerage account in your tax return under the section on foreign affairs and income.

EGYPT

NOTIFICATIONS

Exchange Control Information. If you transfer funds into Egypt in connection with the Units, you are required to transfer the funds through a registered bank in Egypt.

FINLAND

NOTIFICATIONS

Foreign Asset/Account Reporting Information. There are no specific reporting requirements with respect to foreign assets/accounts. However, please note that you must check your pre-completed tax return to confirm that the ownership of Shares and other securities (foreign or domestic) are correctly reported. If you find any errors or omissions, you must make the necessary corrections electronically or by sending specific paper forms to the local tax authorities.

FRANCE

TERMS AND CONDITIONS

Language Consent. By accepting the grant, you confirm having read and understood the Plan and Agreement which were provided in the English language. You accept the terms of these documents accordingly.

Consentement Relatif à la Langue Utilisée. *En acceptant l'attribution, vous confirmez avoir lu et compris le Plan et le Contrat, qui ont été communiqués en langue anglaise. Vous acceptez les termes de ces documents en connaissance de cause.*

NOTIFICATIONS

Foreign Asset/Account Reporting Information. French residents and non-residents must declare to the Customs Authorities the cash and securities they import or export without the use of a financial institution when the value of such cash or securities exceeds a certain threshold (currently €10,000). French residents also must report all foreign bank and brokerage accounts on an annual basis (including accounts opened or closed during the tax year) on Form N° 3916, together with the income tax return. Failure to comply could trigger significant penalties.

GERMANY

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If your acquisition of Shares under the Plan leads to a qualified participation at any point during the calendar year, you will need to report the acquisition

when you file your tax return for the relevant year. A qualified participation is attained only in the unlikely event (i) you own at least 1% of the Company and the value of the Shares acquired exceeds a certain threshold (currently €150,000) or (ii) you hold Shares exceeding 10% of the Company's total Common Stock.

Exchange Control Information. Cross-border payments in excess of a certain threshold (currently €50,000) must be reported monthly to the German Federal Bank (*Bundesbank*). In case of payments in connection with securities (including proceeds realized upon the sale of Shares or the receipt of Dividends or Dividend Equivalents) and/or if the Company withholds Shares with a value in excess of the applicable threshold for any Tax Obligations, the report must be made monthly following the month in which the payment was received and must be filed electronically. The form of report (*Allgemeines Meldeportal Statistik*) can be accessed via the *Bundesbank's* website (www.bundesbank.de) and is available in both German and English. In addition, you may be required to report the acquisition or sale of Shares to the *Bundesbank* if the value of the Shares acquired or sold exceeds the applicable threshold. You are responsible for satisfying any applicable reporting obligation.

GREECE

NOTIFICATIONS

Foreign Asset/Account Reporting Information. The reporting of foreign assets (including Shares and other investments) is your own obligation and takes place through your annual tax return.

HONG KONG

TERMS AND CONDITIONS

Form of Settlement – Units Payable Only in Shares. Notwithstanding any discretion in Section 9.5 of the Plan or anything to the contrary in the Agreement, the Units do not provide any right for you to receive a cash payment and shall be paid in Shares only.

Sale of Shares. Shares received at vesting are accepted as a personal investment. In the event that Shares are issued in respect of the Units within six (6) months of the Grant Date, you agree that you will not offer to the public or otherwise dispose of the Shares prior to the six (6)-month anniversary of the Grant Date.

NOTIFICATIONS

SECURITIES WARNING: *The contents of this document have not been reviewed by any regulatory authority in Hong Kong. You should exercise caution in relation to the offer. If you are in doubt about any of the contents of the Agreement, including this Appendix, or the Plan, you should obtain independent professional advice. The Units and any Shares issued in respect of the Units do not constitute a public offering of securities under Hong Kong law and are available only to members of the Board and Employees. The Agreement, including this Appendix, the Plan and other incidental communication materials have not been prepared in accordance with and are not*

intended to constitute a “prospectus” for a public offering of securities under the applicable securities legislation in Hong Kong. The Units and any documentation related thereto are intended solely for the personal use of each member of the Board and/or Employee and may not be distributed to any other person.

HUNGARY

There are no country-specific provisions.

ICELAND

NOTIFICATIONS

Exchange Control Information. Approval by the Central Bank of Iceland is no longer required to participate in the Plan, regardless of the value of the Shares acquired under the Plan. Despite the recent relaxation of the exchange control requirements, you should consult with your personal advisor to ensure compliance with applicable exchange control regulations in Iceland as such regulations are subject to frequent change. You are responsible for ensuring compliance with all exchange control laws in Iceland.

INDIA

NOTIFICATIONS

Exchange Control Information. You understand that you must repatriate any cash Dividends paid on Shares acquired under the Plan to India or any Dividend Equivalents paid in cash, as well as any proceeds from the sale of Shares acquired under the Plan within such time as may be required under applicable Indian exchange control laws, which may be amended from time to time, unless an exemption applies. You will receive a foreign inward remittance certificate (“**FIRC**”) from the bank where you deposit the foreign currency, and you must maintain the FIRC as proof of repatriation of funds in the event that the Reserve Bank of India or your Employer requests proof of repatriation. It is your responsibility to comply with these requirements. Neither the Company nor the Employer will be liable for any fines or penalties resulting from your failure to comply with any applicable laws. You may be required to provide information regarding funds received from participation in the Plan to the Company and/or the Employer to enable them to comply with their filing requirements under exchange control laws in India.

Foreign Asset/Account Reporting Information. You are required to declare foreign bank accounts and any foreign financial assets (including Shares held outside of India) in your annual tax return. It is your responsibility to comply with this reporting obligation, and you should consult your personal tax advisor in this regard.

IRELAND

TERMS AND CONDITIONS

Acknowledgement of Nature of Plan and Units. This provision supplements Section XI of the Agreement:

In accepting this Agreement, you understand and agree that the benefits received under the Plan will not be taken into account for any redundancy or unfair dismissal claim.

ITALY

TERMS AND CONDITIONS

Acknowledgement of Nature of Agreement. In accepting this Agreement, you acknowledge that (1) you have received a copy of the Plan, the Agreement and this Appendix; (2) you have reviewed the applicable documents in their entirety and fully understand the contents thereof; and (3) you accept all provisions of the Plan, the Agreement and this Appendix.

For any Units granted, you further acknowledge that you have read and specifically and explicitly approve, without limitation, the following sections of the Agreement: Section I; Section II; Section III; Section VIII; Section X; Section XI; Section XVI; Section XVII; and the Data Privacy Notice for All European Economic Area (“EEA”) / European Union (“EU”) Jurisdictions, United Kingdom and Switzerland in this Appendix.

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Italian residents who, at any time during the fiscal year, hold foreign financial assets (including cash and Shares) which may generate income taxable in Italy are required to report these assets on their annual tax returns (UNICO Form, RW Schedule) for the year during which the assets are held, or on a special form if no tax return is due. These reporting obligations will also apply to Italian residents who are the beneficial owners of foreign financial assets under Italian money laundering provisions.

Foreign Financial Assets Tax. The fair market value of any Shares held outside of Italy is subject to a foreign assets tax at a flat rate. The fair market value is considered to be the value of the Shares on the Nasdaq Global Select Market on December 31 of the applicable year in which you held the Shares (or when the Shares are acquired during the course of the year, the tax is levied in proportion to the actual days of holding over the calendar year). No tax payment duties arise if the amount of the foreign financial assets tax calculated on all financial assets held abroad does not exceed a certain threshold. You should consult with your personal tax advisor about the foreign financial assets tax.

JAPAN

NOTIFICATIONS

Foreign Asset/Account Reporting Information. You will be required to report to the Japanese tax authorities details of any assets held outside of Japan as of December 31st (including any

Shares acquired under the Plan) to the extent such assets have a total net fair market value exceeding a certain threshold (currently ¥50,000,000). Such report will be due by June 30 each year. You should consult with your personal tax advisor as to whether the reporting obligation applies to you and whether you will be required to include in the report details of any Shares or cash that you hold.

KOREA

NOTIFICATIONS

Exchange Control Information. Korean residents who sell Shares acquired under the Plan and/or receive cash Dividends on the Shares must file a report with a Korean foreign exchange bank if the proceeds exceed a certain threshold (currently US\$5,000 per transaction) and are deposited into a non-Korean bank account. The report is not required if proceeds are deposited into a non-Korean brokerage account. It is your responsibility to ensure compliance with any applicable exchange control reporting obligations.

Foreign Asset/Account Reporting Information. You are required to declare all foreign financial accounts (*e.g.* non-Korean bank accounts, brokerage accounts holding Shares, etc.) to the Korean tax authority and file a report regarding such accounts if the monthly balance of such accounts exceeds a certain threshold. It is your responsibility to comply with this reporting obligation, and you should consult your personal tax advisor to ensure compliance with this requirement.

LITHUANIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If you (i) hold certain job positions established by the law or (ii) donate to political parties or political campaigners, you must file an Annual Asset Return of the Individual (Family) in Form No. FR0001 with respect to assets held outside of Lithuania (*e.g.*, Shares). If you open an account in a foreign financial institution and annual turnover in the account exceeds a certain threshold (currently €15,000), you must file a foreign account report.

MEXICO

TERMS AND CONDITIONS

Acknowledgement of the Agreement. In accepting the Award granted hereunder, you acknowledge that you have received a copy of the Plan, have reviewed the Plan and the Agreement, including this Appendix, in their entirety and fully understand and accept all provisions of the Plan and the Agreement, including this Appendix. You further acknowledge that you have read and specifically and expressly approve the terms and conditions of Section XI of the Agreement, in which the following is clearly described and established:

- (1) Your participation in the Plan does not constitute an acquired right.

- (2) The Plan and your participation in the Plan are offered by Amgen Inc. on a wholly discretionary basis.
- (3) Your participation in the Plan is voluntary.
- (4) Amgen Inc. and its Affiliates are not responsible for any decrease in the value of the Units granted and/or Shares issued under the Plan.

Labor Law Acknowledgement and Policy Statement. In accepting any Award granted hereunder, you expressly recognize that Amgen Inc., with registered offices at One Amgen Center Drive, Thousand Oaks, California 91320, U.S.A., is solely responsible for the administration of the Plan and that your participation in the Plan and acquisition of Shares do not constitute an employment relationship between you and Amgen Inc. since you are participating in the Plan on a wholly commercial basis and your sole employer is Amgen Mexico S.A. de C.V. ("Amgen-Mexico"). Based on the foregoing, you expressly recognize that the Plan and the benefits that you may derive from participation in the Plan do not establish any rights between you and your Employer, Amgen-Mexico, and do not form part of the employment conditions and/or benefits provided by Amgen-Mexico and any modification of the Plan or its termination shall not constitute a change or impairment of the terms and conditions of your employment.

You further understand that your participation in the Plan is as a result of a unilateral and discretionary decision of Amgen Inc.; therefore, Amgen Inc. reserves the absolute right to amend and/or discontinue your participation in the Plan at any time without any liability to you.

Finally, you hereby declare that you do not reserve to yourself any action or right to bring any claim against Amgen Inc. for any compensation or damages regarding any provision of the Plan or the benefits derived under the Plan, and you therefore grant a full and broad release to Amgen Inc., its Affiliates, stockholders, officers, agents or legal representatives with respect to any claim that may arise.

Spanish Translation

Reconocimiento del Otorgamiento. Al aceptar cualquier Otorgamiento bajo el presente documento, usted reconoce que ha recibido una copia del Plan, que ha revisado el mismo en su totalidad, así como también el Acuerdo de Opción, el Acuerdo, incluyendo este Apéndice, además que comprende y está de acuerdo con todas las disposiciones tanto del Plan y del Otorgamiento, incluyendo este Apéndice. Asimismo, usted reconoce que ha leído y manifiesta específicamente y expresamente la conformidad con los términos y condiciones establecidos en la Sección XI del Acuerdo, en los que se establece y describe claramente que:

- (1) Su participación en el Plan de ninguna manera constituye un derecho adquirido.
- (2) El Plan y su participación en el mismo son ofrecidos por Amgen Inc. de forma completamente discrecional.

- (3) Su participación en el Plan es voluntaria.
- (4) Amgen Inc. y sus Afiliados no son responsables de ninguna disminución en el valor de Unidades o de las Acciones Comunes emitidas mediante el Plan.

Reconocimiento de la Ley Laboral y Declaración de Política. Al aceptar cualquier Otorgamiento de Acciones bajo el presente, usted reconoce expresamente que Amgen Inc., con oficinas registradas localizadas en One Amgen Center Drive, Thousand Oaks, California 91320, U.S.A., es la única responsable de la administración del Plan y que su participación en el mismo y la adquisición de Acciones Comunes no constituyen de ninguna manera una relación laboral entre usted y Amgen Inc., debido a que su participación en el Plan es únicamente una relación comercial y que su único empleador es Amgen Mexico S.A. de C.V. ("Amgen-México"). Derivado de lo anterior, usted reconoce expresamente que el Plan y los beneficios a su favor que pudieran derivar de la participación en el mismo, no establecen ningún derecho entre usted y su empleador, Amgen – México, y no forman parte de las condiciones laborales y/o los beneficios otorgados por Amgen – México, y cualquier modificación del Plan o la terminación del mismo no constituirá un cambio o desmejora de los términos y condiciones de su trabajo.

Asimismo, usted entiende que su participación en el Plan es resultado de la decisión unilateral y discrecional de Amgen Inc., por lo tanto, Amgen Inc. se reserva el derecho absoluto de modificar y/o discontinuar su participación en el Plan en cualquier momento y sin ninguna responsabilidad para usted.

Finalmente, usted manifiesta que no se reserva ninguna acción o derecho que origine una demanda en contra de Amgen Inc., por cualquier compensación o daños y perjuicios, en relación con cualquier disposición del Plan o de los beneficios derivados del mismo, y en consecuencia usted exime amplia y completamente a Amgen Inc. de toda responsabilidad, como así también a sus Afiliadas, accionistas, directores, agentes o representantes legales con respecto a cualquier demanda que pudiera surgir.

NOTIFICATIONS

Securities Law Information. The Units and the Shares offered under the Plan have not been registered with the National Register of Securities maintained by the Mexican National Banking and Securities Commission and cannot be offered or sold publicly in Mexico. In addition, the Plan, the Agreement and any other document relating to the Units may not be publicly distributed in Mexico. These materials are addressed to you only because of your existing relationship with the Company and your Employer and these materials should not be reproduced or copied in any form. The offer contained in these materials does not constitute a public offering of securities but rather constitutes a private placement of securities addressed specifically to individuals who are present employees of Amgen-Mexico made in accordance with the provisions of the Mexican Securities Market Law, and any rights under such offering shall not be assigned or transferred.

NETHERLANDS

NOTIFICATIONS

Securities Law Information.

**Attention! This investment falls outside AFM supervision.
No prospectus required for this activity.**



NORWAY

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Norwegian residents may be subject to foreign asset reporting as part of their ordinary tax return. Norwegian banks, financial institutions, limited companies etc. must report certain information to the Tax Administration. Such information may then be pre-completed in a Norwegian resident's tax return. However, if the resident has traded, or is the owner of, financial instruments (*e.g.*, Shares) not pre-completed in the tax return, the Norwegian resident must enter this information in Form RF-1159, which is an appendix to the tax return.

Exchange Control Information. In general, Norwegian residents should not be subject to any foreign exchange requirements in connection with their acquisition or sale of Shares under the Plan, except normal reporting requirements to the Norwegian Currency Registry. If any transfer of funds into or out of Norway is made through a Norwegian bank, the bank will make the registration.

POLAND

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Polish residents holding foreign securities (including Shares) and maintaining accounts abroad must file reports with the National Bank of Poland if the aggregate value of cash and securities held in such foreign accounts exceeds a certain threshold (currently PLN 7 million). If required, the reports are due on a quarterly basis by the 20th day following the end of each quarter and must be filed on special forms available on the website of the National Bank of Poland.

Exchange Control Information. In addition, Polish residents are required to transfer funds through a bank account in Poland if the transferred amount in any single transaction exceeds a specified threshold (currently €15,000 (or PLN 15,000 if such transfer of funds is associated with the business activity of a consultant)). You must store all documents connected with any foreign exchange transactions you

engage in for a period of five (5) years from the end of the year when such transactions were made. Penalties may apply for failure to comply with exchange control requirements.

PORTUGAL

TERMS AND CONDITIONS

Consent to Receive Information in English. You hereby expressly declare that you have full knowledge of the English language and have read, understood and fully accepted and agreed with the terms and conditions established in the Plan and Agreement.

Conhecimento da Língua. *Por meio do presente, eu declaro expressamente que tem pleno conhecimento da língua inglesa e que li, compreendi e livremente aceitei e concordei com os termos e condições estabelecidas no Plano e no Acordo.*

PUERTO RICO

There are no country-specific provisions.

ROMANIA

NOTIFICATIONS

Exchange Control Information. Certain transfers of funds may need to be reported to the National Office for Prevention and Control of Money Laundering on specific forms by the relevant bank or financial institution. If you deposit proceeds from the sale of Shares or the receipt of Dividends or Dividend Equivalents in a bank account in Romania, you may be required to provide the Romanian bank assisting with the transaction with appropriate documentation explaining the source of the income. You should consult with a legal advisor to determine whether you will be required to submit such documentation to the Romanian bank.

RUSSIA

NOTIFICATIONS

Foreign Account Report Information. Russian residents maintaining accounts abroad must disclose such accounts with the Russian tax authorities, as follows: (i) file notifications upon opening (or closing or changing bank details) within 30 days, and (ii) file reports on the movement of cash or financial assets on such accounts by June 1 of the following year. Such notifications and reports must be filed on

special forms available on the website of the Russian tax authorities. Russian residents who are absent from Russia for more than 183 days in a year may be relieved from this reporting.

TERMS AND CONDITIONS

Securities Law Information. The sale of Shares acquired under the Plan is not permitted within Russia. The Shares are currently traded on the Nasdaq Global Select Market in the U.S. under the ticker symbol “AMGN,” therefore, the Shares may be sold on this exchange. You understand that the exchange control rules and regulations in Russia, legal restrictions impacting your participation in the Plan, are subject to frequent change. You should consult with your personal legal advisor to determine the applicability of any requirements or restrictions applicable to any Shares or cash received in connection with the Plan.”

You understand that the exchange control rules and regulations in Russia and the legal restrictions impacting your participation in the Plan are subject to frequent change. You should consult with your personal legal advisor to determine the applicability of any requirements or restrictions applicable to any Shares or cash received in connection with the Plan.

SINGAPORE

TERMS AND CONDITIONS

Restriction on Sale and Transferability. You hereby agree that any Shares acquired pursuant to the Units will not be offered for sale in Singapore prior to the six (6)-month anniversary of the Grant Date, unless such sale or offer is made pursuant to one or more exemptions under Part XIII Division 1 Subdivision (4) (other than section 280) of the Securities and Futures Act (Chap. 289, 2006 Ed.) (“SFA”), or pursuant to, and in accordance with the conditions of, any other applicable provisions of the SFA.

NOTIFICATIONS

Securities Law Information. The grant of the Units is being made pursuant to the “Qualifying Person” exemption under section 273(1)(f) of the SFA, on which basis it is exempt from the prospectus and registration requirements under the SFA, and is not made with a view to the Units being subsequently offered for sale to any other party. The Plan has not been, and will not be, lodged or registered as a prospectus with the Monetary Authority of Singapore.

Director Notification Requirement. Directors (including alternate, substitute, associate and shadow directors) of a Singapore Affiliate are subject to certain notification requirements under the Singapore Companies Act, regardless of whether they are resident or employed in Singapore. Directors of a Singapore Affiliate must notify the Singapore Affiliate in writing of an interest (e.g., Units, Shares, etc.) in the Company or any related company within two (2) business days of (i) its acquisition or disposal, (ii) any change in a previously disclosed interest (e.g., when the Shares are sold), or (iii) becoming a director.

SLOVAK REPUBLIC

There are no country-specific provisions.

SLOVENIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Slovenian residents may be required to report the opening of bank and/or brokerage accounts to tax authorities within eight (8) days of opening such account. You should consult with your personal tax advisor to determine whether this requirement will be applicable to any accounts opened in connection with your participation in the Plan (*e.g.*, your brokerage account with the Company's designated broker).

SPAIN

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section XI of the Agreement:

By accepting the Units granted hereunder, you consent to participation in the Plan and acknowledge that you have received a copy of the Plan.

You understand that the Company has unilaterally, gratuitously and in its sole discretion decided to grant any Units under the Plan to individuals who may be members of the Board or Employees of the Company or its Affiliates throughout the world. The decision is a limited decision, which is entered into upon the express assumption and condition that (i) any Units granted will not economically or otherwise bind the Company or any of its Affiliates on an ongoing basis, other than as expressly set forth in the Agreement, including this Appendix; (ii) the Units granted hereunder and any Shares acquired under the Plan shall not become a part of any employment contract (either with the Company or any of its Affiliates) and shall not be considered a mandatory benefit, salary for any purposes (including severance compensation) or any other right whatsoever; and (iii) any unvested Units will be forfeited immediately upon termination of your employment for any reason, as detailed below. Further, you understand and freely accept that there is no guarantee that any benefit whatsoever shall arise from any gratuitous and discretionary grant of Units since the future value of the Units and the underlying Shares is unknown and unpredictable. In addition, you understand that any Units granted hereunder would not be made but for the assumptions and conditions referred to above; thus, you understand, acknowledge and freely accept that, should any or all of the assumptions be mistaken or should any of the conditions not be met for any reason, then any grant of Units or right to Units shall be null and void.

Further, the vesting of the Units is expressly conditioned on your continued and active rendering of service, such that if your employment terminates for any reason whatsoever, the Units may

cease vesting immediately, in whole or in part, effective on the date of your termination of employment (unless otherwise specifically provided in Section I of the Agreement). This will be the case, for example, even if (1) you are considered to be unfairly dismissed without good cause (*i.e.*, subject to a “despido improcedente”); (2) you are dismissed for disciplinary or objective reasons or due to a collective dismissal; (3) you terminate service due to a change of work location, duties or any other employment or contractual condition; (4) you terminate service due to a unilateral breach of contract by the Company or an Affiliate; or (5) your employment terminates for any other reason whatsoever. Consequently, upon termination of your employment for any of the above reasons, you may automatically lose any rights to Units that were not vested on the date of your termination of employment, as described in the Plan and the Agreement.

You acknowledge that you have read and specifically accept the conditions referred to in Section I of the Agreement.

NOTIFICATIONS

Securities Law Information. No “offer of securities to the public,” as defined under Spanish law, has taken place or will take place in the Spanish territory. The Agreement (including this Appendix) has not been nor will it be registered with the *Comisión Nacional del Mercado de Valores*, and does not constitute a public offering prospectus.

Exchange Control Information. You will be required to declare electronically to the Bank of Spain any security accounts (including brokerage accounts held abroad), as well as the securities (including Shares acquired under the Plan) held in such accounts if the value of the transactions for all such accounts during the prior year or the balances of such accounts as of December 31 of the prior year exceeds a certain threshold (currently €1,000,000). Different thresholds and deadlines to file this declaration apply. However, if neither such transactions during the immediately preceding year nor the balances / positions as of December 31 exceed the applicable threshold, no such declaration must be filed unless expressly required by the Bank of Spain. If any of such thresholds were exceeded during the current year, you may be required to file the relevant declaration corresponding to the prior year, however, a summarized form of declaration may be available.

Foreign Asset/Account Reporting Information. To the extent that you hold Shares and/or have bank accounts outside of Spain with a value in excess of €50,000 (for each type of asset) as of December 31 each year, you will be required to report information on such assets in your tax return (tax form 720) for such year. After such Shares and/or accounts are initially reported, the reporting obligation will apply for subsequent years only if the value of any previously-reported Shares or accounts increases by more than €20,000 (for each type of asset) as of each subsequent December 31, or if you sell or otherwise dispose of previously-reported Shares or accounts. If the value of such Shares and/or accounts as of December 31 does not exceed €50,000, a summarized form of declaration may be presented.

SWEDEN

TERMS AND CONDITIONS

Authorization to Withhold. This provision supplements Section III of the Agreement:

Without limiting the Company's and the Employer's authority to satisfy their withholding obligations for Tax Obligations as set forth in the Agreement, in accepting the Units, you authorize the Company to withhold Shares or to sell Shares otherwise issuable to you upon vesting or settlement to satisfy Tax Obligations, regardless of whether the Company and/or Employer have an obligation to withhold such Tax Obligations, provided that such withholding would not, in the Company's determination, result in adverse accounting consequences to the Company.

SWITZERLAND

NOTIFICATIONS

Securities Law Information. Neither this document nor any other materials relating to the Awards (i) constitutes a prospectus according to articles 35 et seq. of the Swiss Federal Act on Financial Services ("FinSA"), (ii) may be publicly distributed or otherwise made publicly available in Switzerland to any person other than an employee of the Company or one of its Subsidiaries or (iii) has been or will be filed with, approved or supervised by any Swiss reviewing body according to article 51 of FinSA or any Swiss regulatory authority, including the Swiss Financial Market Supervisory Authority.

TAIWAN

NOTIFICATIONS

Securities Law Information. Participation in the Plan is available only for select Board members and Employees of the Company and its Affiliates. The grant of Units and participation in the Plan is not a public offer of securities by a Taiwanese company.

Exchange Control Information. You may acquire and remit foreign currency (including proceeds from the sale of Shares or the receipt of Dividends or Dividend Equivalents) up to US\$10,000,000 per year without justification. If the transaction amount is TWD500,000 or more in a single transaction, you must submit a Foreign Exchange Transaction Form. If the transaction amount is US\$500,000 or more in a single transaction, you must also provide supporting documentation to the satisfaction of the remitting bank.

THAILAND

NOTIFICATIONS

Exchange Control Information. If proceeds from the sale of Shares or the receipt of any Dividends or Dividend Equivalents exceed a certain threshold (currently US\$1,000,000), you must (i) immediately repatriate such funds to Thailand and (ii) report the inward remittance to the Bank of Thailand on a Foreign Exchange Transaction Form. In addition, within three hundred and sixty (360) days of repatriation, you must either convert any funds repatriated to Thailand to Thai Baht or deposit the funds in a foreign exchange account with a Thai commercial bank. Any such commercial bank must be duly authorized by the Bank of Thailand to engage in the purchase, exchange and withdrawal of foreign currency. The Employee is responsible for ensuring compliance with all exchange control laws in Thailand.

TÜRKİYE

NOTIFICATIONS

Securities Law Information. The sale of Shares acquired under the Plan is not permitted within Türkiye. The sale of Shares acquired under the Plan must occur outside of Türkiye. The Shares are currently traded on the Nasdaq Global Select Market in the U.S. under the ticker symbol “AMGN” and Shares may be sold on this exchange.

Exchange Control Information. You may be required to engage a Turkish financial intermediary to assist with the sale of Shares acquired under the Plan. To the extent a Turkish financial intermediary is required in connection with the sale of any Shares acquired under the Plan, you are solely responsible for engaging such Turkish financial intermediary. You should consult your personal legal advisor prior to the sale of Shares to ensure compliance with the current requirements.

UNITED ARAB EMIRATES

NOTIFICATIONS

Securities Law Information. Units under the Plan are granted only to select Board members and Employees of the Company and its Affiliates and are for the purpose of providing equity incentives. The Plan and the Agreement are intended for distribution only to such Board members and Employees and must not be delivered to, or relied on by, any other person. You should conduct your own due diligence on the Units offered pursuant to this Agreement. If you do not understand the contents of the Plan and/or the Agreement, you should consult an authorized financial adviser. The Emirates Securities and Commodities Authority and the Dubai Financial Services Authority have no responsibility for reviewing or verifying any documents in connection with the Plan. Further, the Ministry of the Economy and the Dubai Department of Economic Development have not approved the Plan or the Agreement nor taken steps to verify the information set out therein, and have no responsibility for such documents.

UNITED KINGDOM

TERMS AND CONDITIONS

Tax Withholding. This provision supplements Section III of the Agreement:

Without limitation to Section III of the Agreement, you agree that you are liable for all Tax Obligations and hereby covenant to pay all such Tax Obligations as and when requested by the Company or your Employer or by HM Revenue and Customs (“**HMRC**”) (or any other tax authority or any other relevant authority). You also agree to indemnify and keep indemnified the Company and your Employer against any taxes that they are required to pay or withhold or have paid or will pay to HMRC (or any other tax authority or any other relevant authority) on your behalf.

Notwithstanding the foregoing, if you are an executive officer or director (as within the meaning of Section 13(k) of the Exchange Act, as amended from time to time), you understand that you may not be able to indemnify the Company or your Employer for the amount of income tax not collected from or paid by you, as it may be considered a loan. In the event that you are an executive officer or director and income tax is not collected from you within ninety (90) days after the end of the tax year in which the Taxable Event occurs, the amount of any uncollected income tax may constitute an additional benefit to you on which additional income tax and national insurance contributions (“**NICs**”) may be payable. You acknowledge that you are responsible for reporting and paying any income tax due on this additional benefit directly to HMRC under the self-assessment regime and for paying your Employer for the amount of any NICs due on this additional benefit, which the Company or your Employer may obtain from you by any of the means set forth in Section III of the Agreement.

If the maximum applicable withholding rate is used, any over-withheld amount may be credited to you by the Company or your Employer (with no entitlement to the Common Stock equivalent) or if not so credited, you may seek a refund from the local tax authorities.

Joint Election. If you are a resident of the United Kingdom between the Grant Date and the vesting of the Units, as a condition of the Units granted hereunder, you agree to accept any liability for secondary Class 1 National Insurance Contributions (the “**Employer NICs**”), which may be payable by the Company or your Employer with respect to the Units and/or payment of the Units and issuance of Shares pursuant to the Units, the assignment or release of the Units for consideration, or the receipt of any other benefit in connection with the Units.

Without limitation to the foregoing, you agree to make an election (the “**Election**”), in the form specified and/or approved for such election by HMRC, that the liability for your Employer NICs payments on any such gains shall be transferred to you to the fullest extent permitted by law. You further agree to execute such other elections as may be required between you and any successor to the Company and/or your Employer. You hereby authorize the Company and your Employer to withhold such Employer NICs by any of the means set forth in Section III of the Agreement.

Failure by you to enter into an Election, withdrawal of approval of the Election by HMRC or a joint revocation of the Election by you and the Company or your Employer, as applicable, shall be grounds for the forfeiture and cancellation of the Units, without any liability to the Company or your Employer.

UNITED STATES

TERMS AND CONDITIONS

Termination of Employment. The following provision replaces Section I(i) of the Agreement:

(i) “termination of your active employment” shall mean the last date that you are either an active employee of the Company or an Affiliate or actively engaged as a Director of the Company or an Affiliate; in the event of termination of your employment (whether or not in breach of local labor laws), your right to receive Units and vest under the Plan, if any, will terminate effective as of the date that you are no longer actively employed; provided, however, that such right will be extended by any notice period mandated by law (e.g., the Worker Adjustment and Retraining Notification Act (“WARN Act”) notice period or similar periods pursuant to local law) and any paid administrative leave (as applicable), unless the Company shall provide you with written notice otherwise before the commencement of such notice period or leave; *provided further*, that notwithstanding the effect of any such extension, in no event will the Units be paid later than the 90th day following your termination of employment;

Form of Award Notice

[The information set forth in this Award Notice will be contained on the related pages on Merrill Lynch Benefits Website (or the website of any successor company to Merrill Lynch Bank & Trust Co., FSB). This Award Notice shall be replaced by the equivalent pages on such website. References to Award Notice in this Agreement shall then refer to the equivalent pages on such website.]

This notice of Award (the “Award Notice”) sets forth certain details relating to the grant by the Company to you of the Award identified below, pursuant to the Plan. The terms of this Award Notice are incorporated into the Performance Unit Agreement (the “Agreement”) that accompanies this Award Notice and made part of the Agreement. Capitalized terms used in this Award Notice that are not otherwise defined in this Award Notice have the meanings given to such terms in the Agreement.

Employee:

Employee ID:

Address:

Award Type:

Grant ID:

Plan: Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan, as amended and/or restated from time to time

Program Amgen Inc. 2009 Performance Award Program, as amended and/or restated from time to time

Grant Date:

Number of Shares:

Number of

Performance Units:

Performance Period: The Performance Period beginning on and ending on .

Resolutions: The Resolutions of the Compensation and Management Development Committee of the Board of Directors of Amgen Inc. establishing the performance goals and Performance Period applicable to this Award.

Vesting Date: Means the vesting date indicated in the Vesting Schedule

Vesting Schedule: Means the schedule of vesting set forth under Vesting Details

Vesting Details: Means the presentation (tabular or otherwise) of the Vesting Date and the quantity of Shares vesting.

IMPORTANT NOTICE REGARDING ACCEPTANCE OF THE AWARD AND THE REQUIREMENT TO OPEN A BROKERAGE ACCOUNT¹:

¹ This provision is only for use on the form of grant used for the U.S. and Puerto Rico.

RESIDENTS OF THE U.S. AND PUERTO RICO: Please read this Award Notice, the Plan and the Agreement (collectively, the “Grant Documents”) carefully. If you, as a resident of the U.S. or Puerto Rico, do **not** wish to receive this Award and/or you do **not** consent and agree to the terms and conditions on which this Award is offered, as set forth in the Grant Documents, then you must reject the Award by contacting the Merrill Lynch call center at +1 (800) 97AMGEN (+1 (800) 972-6436) within the U.S., Puerto Rico and Canada or +1 (609) 818-8910 from all other countries (Merrill Lynch will accept the charges for your call) no later than the forty-fifth calendar day following the day on which this Award Notice is made available to you, in which case the Award will be cancelled. For the purpose of determining the forty-five calendar days, Day 1 will be the day **immediately** following the day on which this Award Notice is made available to you. Your failure to notify the Company of your rejection of the Award or your refusal of, or disagreement with, all terms and conditions of the Award, as set forth in the Grant Documents, within this specified period will constitute your acceptance of the Award and your agreement with all terms and conditions of the Award, as set forth in the Grant Documents. If you agree to the terms and conditions of your grant and you desire to accept it, then no further action is needed on your part to accept the grant. However, you must still open a brokerage account as directed by the Company, by 1:00 pm Pacific Time on or before the date that is 11 months after the Grant Date specified in this Award Notice. This step is necessary to process transactions related to your equity grant. **If you do not open a brokerage account by this deadline, your grant will be cancelled.**

PERFORMANCE UNIT AGREEMENT

THE SPECIFIC TERMS OF YOUR GRANT OF PERFORMANCE UNITS ARE FOUND IN THE PAGES RELATING TO THE GRANT OF PERFORMANCE UNITS FOUND ON MERRILL LYNCH BENEFITS WEBSITE (OR THE WEBSITE OF ANY SUCCESSOR COMPANY TO MERRILL LYNCH BANK & TRUST CO., FSB) (THE “AWARD NOTICE”) WHICH ACCOMPANIES THIS DOCUMENT. THE TERMS OF THE AWARD NOTICE ARE INCORPORATED INTO THIS PERFORMANCE UNIT AGREEMENT.

On the Grant Date specified in the Award Notice, Amgen Inc., a Delaware corporation (the “Company”), has granted to you, the grantee named in the Award Notice, an award under the Amgen Inc. Second Amended and Restated 2009 Equity Incentive Plan, as amended and/or restated from time to time (the “Plan”), for the Number of Performance Units (the “Performance Units”) specified in the Award Notice on the terms and conditions set forth in this Performance Unit Agreement, the Award Notice (and any applicable additional terms and conditions for your country set forth in the attached Appendix A (as described in greater detail in Section XIV below)) (collectively, this “Agreement”), the Plan, the Amgen Inc. 2009 Performance Award Program, as amended and/or restated from time to time (the “Program”) and the Resolutions (as defined in the Award Notice). Capitalized terms not defined herein shall have the meanings assigned to such terms in the Program.

I. Performance Period. The Performance Period shall have the meaning set forth in the Award Notice.

II. Value of Performance Units. The value of each Performance Unit is equal to a share of Common Stock.

III. Performance Goals. An amount of the Performance Units up to the maximum amount specified in the Resolutions shall be earned, depending on the extent to which the Company achieves the performance goals established by the Committee pursuant to the Resolutions. The Performance Units earned shall be calculated in accordance with the Resolutions and the Program.

IV. Form and Timing of Settlement.

(a) General. Subject to Section XIII and the satisfaction of applicable Tax Obligations and similar obligations as provided in Section V, and except as set forth in the Program, any Performance Units earned pursuant to Section III above shall be settled by the Company delivering to you a number of Shares equal to the number of Shares covered by the earned Performance Units or in a lump sum in cash with a value equal to the Fair Market Value of the number of Shares subject to the earned Performance Units as of the last day of the Performance Period (without interest thereon), or in a combination of Shares and cash, as determined by the Administrator at any time prior to settlement and in its discretion, as soon as practicable, and in any event within 90 days, after the last day of the Performance Period. Shares issued in respect of a Performance Unit shall be deemed to be issued in consideration of past services actually rendered by you to the Company or an Affiliate or for its benefit for which you have not

previously been compensated or for future services to be rendered, as the case may be, which the Company deems to have a value at least equal to the aggregate par value thereof.

- (b) Voluntary Retirement. In the event that your employment with the Company or an Affiliate is terminated prior to the last business day of the Performance Period by reason of your Voluntary Retirement and you are Retirement-Eligible on the date of such termination, the full or prorated amount of your Award, if any, applicable to the Performance Period shall be paid in accordance with paragraph (a) of this Section IV. For purposes of the foregoing, the amount of your Award (rounded down to the nearest whole number) shall be determined based on the Company's performance as compared to the Performance Goals for the Performance Period and (i) if the Award was granted with respect to a Performance Period commencing in a calendar year prior to the calendar year in which your Voluntary Retirement occurs, the full amount of the Award is payable, and (ii) if the Award was granted with respect to the Performance Period commencing in the calendar year in which your Voluntary Retirement occurs, the Award otherwise payable is multiplied by a fraction (rounded to two decimal places), the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12). Notwithstanding the foregoing, you shall not be entitled to such full or prorated amount of your Award pursuant to this paragraph (b) unless either you execute and do not revoke a general release and waiver in a form provided by the Company (for the purpose of resolving any potential or actual disputes arising from your employment and the termination of your employment with the Company) (a "Release") and deliver it to the Company no later than the date specified by the Company, or the Company waives such release requirement in writing; *provided, however*, that in no event shall payment of such full or prorated amount of your Award be made later than the specified payment date as set forth in paragraph (a) of this Section IV. This paragraph (b) shall supersede Section 7.1(a) of the Program.
- (c) Death and Disability. In the event that your employment with the Company or an Affiliate is terminated prior to the last business day of the Performance Period by reason of your death or Permanent and Total Disability, the full or prorated amount of your Award, if any, applicable to such Performance Period shall be paid in accordance with the provisions of paragraph (a) of this Section IV. For purposes of the foregoing, the amount of your Award (rounded down to the nearest whole number) shall be determined based on the Company's performance as compared to the Performance Goals for the Performance Period and (i) if the Award was granted with respect to a Performance Period commencing in a calendar year prior to the calendar year in which such termination occurs, the full amount of the Award is payable, and (ii) if the Award was granted with respect to the Performance Period commencing in the calendar year in which such termination occurs, the portion of the Award payable is determined by multiplying the full amount of the Award by a fraction (rounded to two decimal places), the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12). Notwithstanding the foregoing, if your employment is terminated due to your Permanent and Total

Disability, you shall not be entitled to such full or prorated amount of your Award pursuant to this paragraph (c) unless either you execute and do not revoke a Release and deliver it to the Company no later than the date specified by the Company, or the Company waives such release requirement in writing; *provided, however*, that in no event shall payment of such full or prorated amount of your Award be made later than the specified payment date as set forth in paragraph (a) of this Section IV. This paragraph (c) shall supersede Section 7.1(b) of the Program.

- (d) *Other*. In the event that your employment with the Company or an Affiliate is terminated prior to the last business day of the Performance Period for any reason other than as specified in paragraphs (b) and (c) above, all of your rights to an Award for the Performance Period shall be forfeited, unless, prior to the payment date set forth in paragraph (a) of this Section IV, the Company, in its sole discretion, makes a written determination to otherwise pay the full or prorated amount of your Award, if any, applicable to the Performance Period, which full or prorated amount shall be paid in accordance with the provisions of paragraph (a) of this Section IV. For purposes of the foregoing, if the payment of your Award is prorated, the amount of your Award (rounded down to the nearest whole number) shall be determined based on the Company's performance as compared to the Performance Goals for the Performance Period and (i) if the Award was granted with respect to a Performance Period commencing in a calendar year prior to the calendar year in which such termination occurs, the full amount of the Award is payable, and (ii) if the Award was granted with respect to the Performance Period commencing in the calendar year in which such termination occurs, the Award otherwise payable is multiplied by a fraction (rounded to two decimal places), the numerator of which is the number of complete months you remained continuously and actively employed by the Company or an Affiliate during the calendar year in which your termination occurs, and the denominator of which is twelve (12). Notwithstanding the foregoing, you shall not be entitled to such full or prorated amount of your Award pursuant to this paragraph (d) unless either you execute and do not revoke a Release and deliver it to the Company no later than the date specified by the Company, or the Company waives such release requirement in writing; *provided, however*, that in no event shall payment of such full or prorated amount of your Award be made later than the specified payment date as set forth in paragraph (a) of this Section IV. This paragraph (d) shall supersede Section 7.1(c) of the Program.

V. Issuance of Shares; Tax Withholding. Regardless of any action the Company or your actual employer (the "Employer") takes with respect to any or all income tax (including federal, state and local taxes), social insurance, payroll tax, fringe benefit tax, payment on account or other tax-related items related to your participation in the Plan and the Program and legally applicable to you (the "Tax Obligations"), you acknowledge that the ultimate liability for all Tax Obligations is and remains your responsibility and may exceed the amount, if any, actually withheld by the Company and/or your Employer. You further acknowledge that the Company and/or your Employer (i) make no representations or undertakings regarding the treatment of any Tax Obligations in connection with any aspect of the Performance Units or the underlying Shares, including the grant of the Performance Units, the vesting of the Performance Units, the conversion of the Performance Units into shares or the receipt of an equivalent cash payment, the subsequent sale of any shares acquired at settlement and the receipt of any

Dividends (as defined in Section VI, below) or Dividend Equivalents; and (ii) do not commit to and are under no obligation to structure the terms of the grant or any aspect of the Performance Units to reduce or eliminate your liability for Tax Obligations or to achieve any particular tax result. Furthermore, if you become subject to tax in more than one jurisdiction, you acknowledge that the Company and/or your Employer (or former employer, as applicable) may be required to withhold or account for Tax Obligations in more than one jurisdiction.

Prior to any relevant taxable or tax withholding event, as applicable, you shall pay or make adequate arrangements satisfactory to the Company or to your Employer (in their sole discretion) to satisfy all Tax Obligations. In this regard, you authorize the Company and/or your Employer, or their respective agents, at their discretion, to satisfy all applicable Tax Obligations by one or a combination of the following:

- (a) withholding from your wages or other cash compensation paid to you by the Company and/or your Employer; or
- (b) withholding from proceeds of the sale of Shares issued upon settlement of the Performance Units, either through your voluntary sale or through a mandatory sale arranged by the Company (on your behalf pursuant to this authorization); or
- (c) withholding in Shares issuable, or cash payable, upon settlement of the Performance Units provided that, if such Shares are withheld, the Company and your Employer shall only withhold an amount of Shares with a fair market value not to exceed the Tax Obligations as determined in the discretion of the Company or your Employer, as applicable.

Depending on the withholding method, the Company may withhold or account for Tax Obligations by considering applicable minimum statutory withholding rates or other applicable withholding rates, including maximum applicable rates. If the Tax Obligations are satisfied by withholding in Shares, for tax purposes you are deemed to have been issued the full number of Shares subject to the earned Performance Units, notwithstanding that a number of Shares is held back and not actually issued to you solely for the purpose of paying the Tax Obligations due as a result of any aspect of your participation in the Plan (any Shares withheld by the Company hereunder shall not be deemed to have been issued by the Company for any purpose under the Plan and shall remain available for issuance thereunder).

Finally, you shall pay to the Company or your Employer any amount of Tax Obligations that the Company or your Employer may be required to withhold or account for as a result of your participation in the Plan and the Program that cannot be or were not satisfied by the means previously described. You agree to take any further actions and to execute any additional documents as may be necessary to effectuate the provisions of this Section V. Notwithstanding Section IV above, the Company may refuse to issue or deliver the Shares or the proceeds of the sale of Shares if you fail to comply with your obligations in connection with the Tax Obligations.

VI. Dividend Equivalents

- (a) Crediting of Dividend Equivalents. Subject to this Section VI, Dividend Equivalents shall be credited on each Performance Unit granted to you under this Agreement in the manner set forth in the remainder of this Section VI. With respect to each Performance Unit covered by the Award, if the Company declares one or more dividends or distributions (each, a "Dividend") on its Common Stock with a record date which occurs during the period commencing on the Grant Date through and including the day immediately preceding the day the Share subject to each Performance Unit is issued to you, whether in the form of cash, Common Stock or other property, then, on the date such Dividend is paid to the Company's stockholders, you shall be credited with an amount equal to the amount or fair market value of such Dividend which would have been payable to you if you held a number of Shares equal to the number of Performance Units granted to you on the Grant Date (including any previously credited Dividends which have been deemed to have been reinvested in Common Stock as provided by the next succeeding sentence), as of each such record date for each such Dividend (not including on any Performance Units which were previously paid or forfeited) as if each such amount had been reinvested in Common Stock as of the date of the payment of such Dividend (such accumulated dividends, the "Target Accumulated Dividends"). Each such Dividend Equivalent shall be deemed to have been reinvested in Common Stock as of the applicable Dividend payment date. Dividend Equivalents shall be payable in full Shares, unless the Administrator determines, at any time prior to payment and in its discretion, that they shall be payable in cash. Dividend Equivalents payable with respect to fractional Shares shall be paid in cash.
- (b) Treatment of Dividend Equivalents. Except as otherwise expressly provided in this Section VI any Dividend Equivalents credited to you shall be subject to all of the provisions of this Agreement which apply to the Performance Units with respect to which they have been credited and shall be payable, if at all, at the time and to the extent that the underlying Performance Unit becomes payable pursuant to paragraph (a) of Section IV above. Dividend Equivalents shall not be payable on any Performance Units that do not vest, or are forfeited, pursuant to the terms of this Agreement. Dividend Equivalent rights and any amounts that may become distributable in respect thereof shall be treated separately from the Performance Units and the rights arising in connection therewith for purposes of the designation of time and form of payments required by Section 409A of the Internal Revenue Code of 1986, as amended (the "Code") (together with any Department of Treasury regulations and other interpretive guidance issued thereunder, including without limitation any such regulations or other guidance that may be issued after the Grant Date, "Section 409A").

VII. Nontransferability. No benefit payable under, or interest in, this Agreement, the Units, or the Shares that may become issuable to you hereunder shall be subject in any manner to anticipation, alienation, sale, transfer, assignment, pledge, encumbrance or charge and any such attempted action shall be void and no such benefit or interest shall be, in any manner, liable for, or subject to, your or your beneficiary's debts, contracts, liabilities or torts; *provided, however*, nothing in this Section VII shall prevent transfer (i) by will or (ii) by applicable laws of descent and distribution.

VIII. No Contract for Employment. This Agreement is not an employment or service contract with the Company or an Affiliate and nothing in this Agreement shall be deemed to create in any way whatsoever any obligation on your part to continue in the employ or service of the Company

or an Affiliate, or of the Company or an Affiliate to continue your employment or service with the Company or an Affiliate.

IX. Nature of Grant. In accepting the grant of Performance Units, you acknowledge, understand and agree that:

- (a) the Plan and the Program are established voluntarily by the Company, are discretionary in nature and may be modified, amended, suspended or terminated by the Company at any time, as provided in the Plan and in the Program;
- (b) the Plan is operated and the Performance Units are granted solely by the Company, and only the Company is a party to this Agreement; accordingly, any rights you may have under this Agreement may be raised only against the Company but not any Affiliate (including, but not limited to, the Employer);
- (c) no Affiliate (including, but not limited to, the Employer) has any obligation to make any payment of any kind to you under this Agreement;
- (d) the grant of the Performance Units is exceptional, voluntary and occasional and does not create any contractual or other right to receive future awards of Performance Units, or benefits in lieu of Performance Units, even if Performance Units have been awarded in the past;
- (e) all decisions with respect to future awards, if any, will be at the sole discretion of the Company;
- (f) your participation in the Plan and the Program is voluntary;
- (g) the grant of Performance Units, the Shares subject to the Performance Units, and the income from and value of same, are not intended to replace any pension rights or compensation;
- (h) neither the grant of Performance Units nor any provision of this Agreement, the Plan, the Program or the policies adopted pursuant to the Plan or Program confer upon you any right with respect to employment or continuation of current employment and shall not interfere with the ability of your Employer to terminate your employment or service relationship (if any) at any time;
- (i) in the event that you are not an employee of the Company or any Affiliate, the Performance Units shall not be interpreted to form an employment contract or relationship with the Company or any Affiliate;
- (j) the future value of the Shares that may be earned upon the end of the Performance Period is unknown, indeterminable, and cannot be predicted with certainty;
- (k) in consideration of the grant of Performance Units hereunder, no claim or entitlement to compensation or damages arises from termination of Performance Units, and no claim or entitlement to compensation or damages shall arise from forfeiture of the Performance Units resulting from (i) the termination of your employment by the Company or an Affiliate (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any) and/or (ii) the application of the Clawback Policy as described in Section XXI below, and you irrevocably release the Company and your Employer from any such claim that may arise; if, notwithstanding the foregoing, any such claim is found by a court

of competent jurisdiction to have arisen, you shall be deemed irrevocably to have waived your entitlement to pursue such claim;

- (l) in the event of termination of your employment (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any), your right to receive Performance Units and receive shares under the Plan and the Program, if any, will terminate effective as of the date that you are no longer actively employed and will not be extended by any notice period (*e.g.*, active employment would not include a period of “garden leave” or similar period mandated under employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any);
- (m) unless otherwise agreed with the Company, the Performance Units, the Shares subject to the Performance Units, and the income from and value of same, are not granted as consideration for, or in connection with, the service you may provide as a director of an Affiliate of the Company;
- (n) except as otherwise provided in this Agreement or the Plan, the Performance Units and the benefits evidenced by this Agreement do not create any entitlement to have the Performance Units or any such benefits transferred to, or assumed by, another company nor to be exchanged, cashed out or substituted for, in connection with any corporate transaction affecting the shares of the Company; and
- (o) the following provisions apply only if you are providing services outside the United States:
 - (A) for employment law purposes outside the United States, the Performance Units, the Shares subject to the Performance Units, and the income from and value of same, are not part of normal or expected *compensation* or salary for any purpose, including but not limited to for purposes of calculating any severance, resignation, termination, redundancy, dismissal, end of service payments, bonuses, holiday pay, long-service awards, pension or retirement benefits or similar mandatory payments; and
 - (B) neither the Company, your Employer nor any Affiliate of the Company shall be liable for any foreign exchange rate fluctuation between your local currency and the United States Dollar that may affect the value of the Performance Units or of any amounts due to you pursuant to the settlement of the Performance Units or the subsequent sale of any Shares acquired upon settlement.

X. No Advice Regarding Grant. The Company is not providing any tax, legal or financial advice, nor is the Company making any recommendations regarding your participation in the Plan and the Program, or your acquisition or sale of the underlying Shares. You should consult with your personal tax, legal and financial advisors regarding your participation in the Plan and the Program before taking any action related thereto.

XI. Notices. Any notices provided for in this Agreement, the Plan or the Program shall be given in writing or electronically and shall be deemed effectively given upon receipt or, in the case of notices delivered by the Company to you, five (5) days after deposit in the United States mail or equivalent foreign postal service, postage prepaid, addressed to you at such address as is currently maintained in the Company’s records or at such other address as you hereafter designate by written notice to the Company Stock Administrator. Such notices may be given using any automated system

for the documentation, granting or settlement of Awards, such as a system using an internet website or interactive voice response, as approved by the Company.

XII. Resolutions, Plan and Program. This Agreement is subject to all of the provisions of the Resolutions, the Plan and the Program and their provisions are hereby made a part of this Agreement and incorporated herein by reference, including, without limitation, the provisions of Article 9 of the Plan (relating to Performance Awards) and Section 13.2 of the Plan (relating to adjustments upon changes in the Common Stock), and is further subject to all interpretations, amendments, rules and regulations which may from time to time be promulgated and adopted pursuant to the Plan. In the event of any conflict between the provisions of this Agreement and those of the Resolutions, the Plan and the Program, the provisions of the Plan shall control. Notwithstanding any provision of this Agreement or the Program to the contrary, any earned Performance Units paid in cash rather than Shares shall not be deemed to have been issued by the Company for any purpose under the Plan.

XIII. Code Section 409A. The time and form of payment of the Performance Units is intended to comply with the requirements of Section 409A and this Agreement shall be interpreted in accordance with Section 409A. Accordingly, no acceleration or deferral of any payment shall be permitted if it would cause the payment of the Performance Units to violate Section 409A. In addition, notwithstanding any provision herein to the contrary, in the event that following the Grant Date, the Committee determines that it may be necessary or appropriate to do so, the Committee may adopt such amendments to the Plan, Program and/or this Agreement or adopt other policies and procedures (including amendments, policies and procedures with retroactive effect), or take any other actions, that the Committee determines are necessary or appropriate to (a) exempt the Plan, Program and/or the Performance Units from the application of Section 409A and/or preserve the intended tax treatment of the benefits provided with respect to this Award, or (b) comply with the requirements of Section 409A; *provided, however*, that this paragraph shall not create an obligation on the part of the Committee to adopt any such amendment, policy or procedure or take any such other action. No payment hereunder shall be made to you during the six (6)-month period following your “separation from service” (within the meaning of Section 409A) to the extent that the Company determines that paying such amount at the time set forth herein would be a prohibited distribution under Section 409A(a)(2)(B)(i). If the payment of any such amounts is delayed as a result of the previous sentence, then within thirty (30) days following the end of such six (6)-month period (or, if earlier, your death), the Company shall pay to you (or to your estate) the cumulative amounts that would have otherwise been payable to you during such period, without interest.

XIV. Provisions Applicable to Participants in Foreign Jurisdictions. Notwithstanding any provision of this Agreement or the Program to the contrary, if you are employed by the Company or an Affiliate in any of the countries identified in the attached Appendix A (which constitutes a part of this Agreement), are subject to the laws of any foreign jurisdiction, or relocate to one of the countries included in the attached Appendix A, your award of Performance Units shall be subject to any additional terms and conditions for such country set forth in Appendix A and to the following additional terms and conditions:

- (a) the terms and conditions of this Agreement, including Appendix A, are deemed modified to the extent necessary or advisable to comply with applicable foreign laws or facilitate the administration of the Plan and the Program;
- (b) if applicable, the effectiveness of your Award is conditioned upon its compliance with any applicable foreign laws, regulations, rules or local governmental regulatory exemption and subject to receipt of any required foreign regulatory approvals;
- (c) to the extent necessary to comply with applicable foreign laws, the payment of any earned Performance Units shall be made in cash or Common Stock, at the Company's election; and
- (d) the Committee may take any other action, before or after an award of Performance Units is made, that it deems necessary or advisable to obtain approval or comply with any necessary local governmental regulatory exemptions or approvals.

Notwithstanding anything to the contrary contained herein, the Company shall not take any actions hereunder, and no Award of Performance Units shall be granted, and no Shares payable with respect to an Award shall be issued, that would violate the Securities Act, the Exchange Act, the Code, or any other securities or tax or other applicable law or regulation, or the rules of any Securities Exchange. Notwithstanding anything to the contrary contained herein, no Shares issuable with respect to an Award shall be issued unless such shares are then registered under the Securities Act, or, if such shares are not then so registered, the Company has determined that such issuance would be exempt from the registration requirements of the Securities Act and that the issuance satisfied all other applicable legal requirements.

XV. Data Privacy. *In order for the Company to facilitate your participation in the Plan and the Program, the Company and your Employer must collect and use personal data about you. In accordance with applicable laws, reasonable security measures will be implemented and maintained to protect the security of your personal data; however, you understand that absolute security cannot be guaranteed.*

You understand that the Company and your Employer may hold certain personal information about you, including your name, home address and telephone number, email address, date of birth, social insurance/security number (to the extent permitted under applicable local law), passport or other identification number, salary, nationality, job title/work history/service periods, residency status, citizenship, tax withholding and payroll data, any shares of stock or directorships held in the Company, details of all equity compensation or any other entitlement to Shares awarded, cancelled, vested, unvested or outstanding in your favor, for the purposes of implementing, administering and managing the Plan and the Program ("personal data").

You authorize the transfer of your personal data to Merrill Lynch Bank & Trust Co., FSB, or any successor thereto, and any other third parties which may assist the Company (presently or in the future) with implementing, administering and managing your participation in the Plan and the Program to receive, possess, use, retain and transfer your personal data, in electronic or other form, for the purpose of implementing, administering and managing your participation in the Plan and the Program, including any requisite transfer of such personal data as may be required to any other broker, escrow agent or other third party with whom the Shares received in settlement of the

Performance Units may be deposited. You understand that such authorized recipients of your personal data may be located in countries that do not provide the same level of data privacy laws and protections as the country in which your personal data originated. Transfers of personal data among Company and its group entities follow applicable laws and our Binding Corporate Rules (BCRs). For more information on Company's BCRs, please visit <http://www.amgen.com/bcr/>. You acknowledge that the collection, use and transfer of your personal data is necessary to facilitate your participation in the Plan, as well as to grant you Performance Units or other equity awards and administer or maintain such awards.

You may correct or update your personal data previously provided to Company, by contacting your local human resources representative. Subject to applicable law, you may have additional rights, including the right to object and/or request destruction of your personal data. To exercise these rights, where applicable, please contact your local human resources representative.

XVI. Language. By electing to accept this Agreement, you acknowledge that you are sufficiently proficient in English or have consulted with an advisor who is sufficiently proficient in English, so as to allow you to understand the terms and conditions of this Agreement. Furthermore, if you have received this Agreement or any other document related to the Plan and/or the Program translated into a language other than English and if the meaning of the translated version is different than the English version, the English version will control, unless otherwise required by applicable law.

XVII. Governing Law and Venue. The terms of this Agreement shall be governed by the laws of the State of Delaware without giving effect to principles of conflicts of laws. For purposes of litigating any dispute that arises hereunder, the parties hereby submit to and consent to the jurisdiction of the State of Delaware, and agree that such litigation shall be conducted in the courts of the State of Delaware, or the federal courts for the United States for the federal district located in the State of Delaware, and no other courts, where this Agreement is made and/or to be performed.

XVIII. Severability. If one or more of the provisions of this Agreement shall be held invalid, illegal or unenforceable in any respect, the validity, legality and enforceability of the remaining provisions shall not in any way be affected or impaired thereby and the invalid, illegal or unenforceable provisions shall be deemed null and void; however, to the extent permissible by law, any provisions which could be deemed null and void shall first be construed, interpreted or revised retroactively to permit this Agreement to be construed so as to foster the intent of this Agreement and the Plan.

XIX. Electronic Delivery and Participation. The Company may, in its sole discretion, decide to deliver any documents related to current or future participation in the Plan and/or the Program (including this Agreement) by electronic means. You hereby consent to receive such documents by electronic delivery and agree to participate in the Plan through an on-line or electronic system established and maintained by the Company or another third party designated by the Company.

XX. Imposition of Other Requirements. The Company reserves the right to impose other requirements on your participation in the Plan and the Program, on the Performance Units and on any Shares acquired under the Plan and the Program, to the extent the Company determines it is necessary

or advisable for legal or administrative reasons, and to require you to sign any additional agreements or undertakings that may be necessary to accomplish the foregoing.

XXI. Compensation Subject to Recovery. The Performance Units subject to and Shares issuable under this Award and all compensation payable with respect to them shall be subject to clawback, recoupment and/or recovery by the Company pursuant to any and all of the Company's policies with respect to the clawback, recoupment or recovery of compensation in effect as of the Grant Date or as may be adopted or maintained by the Company following the Grant Date, including, without limitation, the Company's Policy on Recovery of Erroneously Awarded Compensation (effective October 2, 2023) and Executive Officer Equity Recoupment Policy (effective December 31, 2020), as they shall be in effect and may be amended from time to time (collectively, the "Clawback Policy"), to the maximum extent permitted by applicable law.

XXII. Waiver. You acknowledge that a waiver by the Company of breach of any provision of this Agreement shall not operate or be construed as a waiver of any other provision of this Agreement, or of any subsequent breach by you or any other grantee.

XXIII. Headings. This Agreement's section headings are for convenience only and shall not constitute a part of this Agreement or affect this Agreement's meaning.

Very truly yours,
AMGEN INC.

By: _____

Name:

Title:

APPENDIX A

ADDITIONAL TERMS AND CONDITIONS OF THE AMGEN INC. SECOND AMENDED AND RESTATED 2009 EQUITY INCENTIVE PLAN, AS AMENDED AND/OR RESTATED FROM TIME TO TIME

AWARD OF PERFORMANCE UNITS (BY COUNTRY)

Certain capitalized terms used but not defined in this Appendix A shall have the meanings set forth in the Plan and/or the Agreement to which this Appendix is attached.

TERMS AND CONDITIONS

This Appendix includes additional terms and conditions that govern any Performance Units granted under the Plan if, under applicable law, you are a resident of, are deemed to be a resident of or are working in one of the countries listed below. Furthermore, the additional terms and conditions that govern the Performance Units granted hereunder may apply to you if you transfer employment and/or residency to one of the countries listed below and the Company shall, in its discretion, determine to what extent the terms and conditions contained herein shall apply to you.

NOTIFICATIONS

This Appendix also includes notifications relating to exchange control and other issues of which you should be aware with respect to your participation in the Plan. The information is based on the exchange control, securities and other laws in effect in the countries to which this Appendix refers as of October 2025. Such laws are often complex and change frequently. As a result, the Company strongly recommends that you not rely on the notifications herein as the only source of information relating to the consequences of your participation in the Plan because the information may be outdated when you acquire Shares under the Plan, or when you subsequently sell Shares acquired under the Plan and the Program.

In addition, the notifications are general in nature and may not apply to your particular situation, and the Company is not in a position to assure you of any particular result. Accordingly, you should seek appropriate professional advice as to how the relevant laws in your country may apply to your situation. Finally, if you are a citizen or resident of a country other than the one in which you are currently residing and/or working or are considered a resident of another country for local law purposes, the information contained herein may not be applicable to you or you may be subject to the provisions of one or more jurisdictions.

ALL NON-U.S. JURISDICTIONS

TERMS AND CONDITIONS

Issuance of Shares; Tax Withholding. The following provision supplements Section V. of the Agreement:

In the event the Company withholds or accounts for Tax Obligations by considering maximum applicable rates in your jurisdiction(s), in the event of over-withholding, you may receive a refund of any over-withheld amount in cash and will not be entitled to the equivalent amount in Shares, or if not refunded, you may seek a refund from the local tax authorities. In the event of under-withholding, you may be required to pay any additional Tax Obligations directly to the applicable tax authority or to the Company and/or your Employer.

NOTIFICATIONS

Insider Trading Restrictions/Market Abuse Laws. You may be subject to insider trading restrictions and/or market abuse laws based on the exchange on which the Shares are listed and in applicable jurisdictions including the United States and your country or your broker's country, if different, which may affect your ability to accept, acquire, sell or otherwise dispose of Shares, rights to Shares (e.g., Performance Units) or rights linked to the value of Shares (e.g., Dividend Equivalents) during such times as you are considered to have "inside information" regarding the Company (as defined by the laws in applicable jurisdictions). Local insider trading laws and regulations may prohibit the cancellation or amendment of orders you place before you possessed inside information. Furthermore, you could be prohibited from (i) disclosing the inside information to any third party, which may include fellow employees (other than on a "need to know" basis) and (ii) "tipping" third parties or causing them otherwise to buy or sell securities. Any restrictions under these laws or regulations are separate from and in addition to any restrictions that may be imposed under any applicable Company insider trading policy. You are responsible for ensuring your compliance with any applicable restrictions and you should speak with your personal legal advisor on this matter.

Foreign Asset/Account, Tax Reporting Information. Your country of residence may have certain foreign asset and/or account reporting requirements which may affect your ability to acquire or hold Shares under the Plan or cash received from participating in the Plan (including from any Dividends or Dividend Equivalents received, or sale proceeds arising from the sale of Shares) in a brokerage or bank account outside of your country. You may be required to report such accounts, assets or transactions to the tax or other authorities in your country. You also may be required to repatriate sale proceeds or other funds received as a result of participating in the Plan to your country within a certain time after receipt. You are responsible for ensuring your compliance with such regulations, and you should speak with your personal legal advisor on this matter.

ALL EUROPEAN ECONOMIC AREA (“EEA”) / EUROPEAN UNION (“EU”) JURISDICTIONS, UNITED KINGDOM AND SWITZERLAND

TERMS AND CONDITIONS

Data Privacy Notice. This provision replaces Section XV of the Agreement:

Please refer to the Fair Processing Notice previously provided by your local human resources representative, which notice governs the collection, use and transfer of your personal data necessary for the Company to facilitate your participation in the Plan and the Program. If you have any questions or concerns regarding the Fair Processing Notice, including questions about your rights afforded thereunder, you should contact your local human resources representative or send an email to hrconnect@amgen.com.

For purposes of implementing, administering and managing the Plan, Company and your Employer may hold certain personal data about you, including your name, home address and telephone number, email address, date of birth, social insurance/security number (to the extent permitted under applicable local law), passport or other identification number, salary, nationality, job title/work history/service periods, residency status, citizenship, tax withholding and payroll data, any shares of stock or directorships held in the Company, details of all equity compensation or any other entitlement to Shares awarded, cancelled, vested, unvested or outstanding in your favor (“personal data”).

You authorize the transfer of your personal data to Merrill Lynch Bank & Trust Co., FSB, or any successor thereto, and any other third parties which may assist the Company (presently or in the future) with implementing, administering and managing your participation in the Plan and the Program to receive, possess, use, retain and transfer your personal data, in electronic or other form, for the purpose of implementing, administering and managing your participation in the Plan and the Program, including any requisite transfer of such personal data as may be required to any other broker, escrow agent or other third party with whom the Shares received in settlement of the Performance Units may be deposited.

ARGENTINA

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section IX of the Agreement:

In accepting the grant of Performance Units, you acknowledge, understand and agree that the grant of the Performance Units is made by the Company (not your Employer) in its sole discretion and that the value of the Performance Units or any Shares acquired under the Plan and the Program shall not constitute salary or wages for any purpose under Argentine labor law including, but not limited to, the calculation of (i) any labor benefits including, without limitation, vacation pay, thirteenth salary, compensation in lieu of notice, annual bonus, disability, and leave of absence payments, etc., or (ii) any termination or severance indemnities or similar payments.

NOTIFICATIONS

Securities Law Information. The Performance Units and the underlying Shares have not been and will not be publicly issued, placed, distributed, offered or listed in the Argentine capital markets, and, as a result, have not been and will not be registered with the Argentina Securities Commission (Comisión Nacional de Valores, “CNV”). Neither this nor any other offering material related to the Performance Units nor the underlying Shares may be utilized in connection with any general offering to the public in Argentina. Any Argentine resident who acquires Shares under the Plan does so under their own responsibility under the terms of a private offering to the Argentine resident from outside Argentina. Any Argentine resident who acquires Shares shall not transfer such Shares to any other person within six (6) months of acquiring the Shares, unless the transaction is conducted outside Argentina.

Exchange Control Information. Exchange control regulations in Argentina are subject to frequent change. You should consult with your personal legal advisor regarding any exchange control obligations that you may have prior to receiving proceeds from Dividend Equivalents, the sale of Shares or dividends. You must comply with any and all Argentine currency exchange restrictions, approvals and reporting requirements in connection with your participation in the Plan and the Program.

Foreign Asset/Account Reporting Information. If you are an Argentine resident, you are required to report certain information regarding any Shares you hold as of December 31 each year to the Argentine tax authorities on your annual tax return.

AUSTRALIA

NOTIFICATIONS

Australia Offer Document. This grant of Units is being made under Division 1A, Part 7.12 of the Corporations Act 2001 (Cth).

Tax Information. Subdivision 83A-C of the Income Tax Assessment Act 1997 (Cth) applies to the Performance Units granted under the Plan, such that the Performance Units are intended to be subject to deferred taxation.

Exchange Control Information. If you are an Australian resident, exchange control reporting is required for cash transactions exceeding a certain threshold (currently AUD10,000) and for international fund transfers. If an Australian bank is assisting with the transaction, the bank will file the report on your behalf. If there is no Australian bank involved in the transfer, you will be required to file the report.

AUSTRIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If you are an Austrian resident and you hold Shares acquired under the Plan and the Program outside of Austria, you may be subject to reporting obligations to the Austrian National Bank.

Exchange Control Information. If the value of Shares meets or exceeds a certain threshold (currently, €5,000,000), you must report the securities held on a quarterly basis to the Austrian National Bank as of the last day of the quarter, on or before the 15th day of the month following the end of the calendar quarter. Further, if you hold cash in accounts outside Austria, monthly reporting requirements will apply, as explained in the next paragraph, if the aggregate transaction volume of such cash accounts meets or exceeds a certain threshold.

A separate reporting requirement applies when you sell Shares acquired under the Plan and the Program, receive a cash Dividend paid on such Shares or Dividend Equivalents paid in cash. In that case, there may be exchange control obligations if the cash proceeds are held outside of Austria. If the transaction volume of all cash accounts abroad meets or exceeds a certain threshold (currently €10,000,000), the movements and balances of all accounts must be reported monthly, as of the last day of the month, on or before the 15th day of the following month, on the prescribed forms.

BELGIUM

NOTIFICATIONS

Tax Reporting; Foreign Asset/Account Reporting Information. If you are a Belgian resident, you are required to report any securities (e.g., Shares acquired under the Plan and the Program) held and bank accounts (including brokerage accounts) opened and maintained outside of Belgium on your annual tax return. The first time you report the foreign security and/or bank account on your annual income tax return you will have to provide the National Bank of Belgium Central Contact Point with the account details of any such foreign accounts (including the account number, bank name and country in which such account was opened) in a separate form. This report, as well as information on how to complete it, can be found on the website of the National Bank of Belgium, www.nbb.be, under the *Kredietcentrales / Centrales des crédits* caption.

Stock Exchange Tax Information. A stock exchange tax applies to transactions executed by a Belgian resident through a non-Belgian financial intermediary, such as a U.S. broker. The stock exchange tax likely will apply when Shares acquired under the Plan and the Program are sold. It is your responsibility to comply with this tax obligation and you should consult your personal tax advisor for additional details on your obligations with respect to the stock exchange tax.

Annual Securities Accounts Tax Information. An annual securities accounts tax may be payable if the total value of securities held in a Belgian or foreign securities account (e.g., Shares acquired under the Plan and the Program) exceeds a certain threshold on four reference dates within the relevant reporting period (i.e., December 31, March 31, June 30 and September 30). In such case, the tax will be due on the value of the qualifying securities held in such account. It is your responsibility to comply with this obligation, and you should consult with your personal tax or financial advisor for additional details.

BRAZIL

TERMS AND CONDITIONS

Compliance with Law. By accepting the Performance Units, you acknowledge that you agree to comply with applicable Brazilian laws and pay any and all applicable taxes associated with the vesting of the Performance Units, the sale of Shares acquired under the Plan and the Program, the payment of Dividends on such Shares and the receipt of any Dividend Equivalents paid in cash.

Nature of Grant. This provision supplements Section IX of the Agreement:

In accepting the grant of Performance Units, you acknowledge (i) that you are making an investment decision, (ii) that the Shares will be issued to you only if the vesting conditions are met and any necessary services are rendered by you during the vesting period set forth in the Vesting Schedule, and (iii) that the value of the underlying Shares is not fixed and may increase or decrease in value over the vesting period without compensation to you.

NOTIFICATIONS

Exchange Control Information. If you are resident or domiciled in Brazil, you will be required to submit annually a declaration of assets and rights held outside of Brazil to the Central Bank of Brazil if the aggregate value of such assets and rights on December 31 of each year exceeds US\$1,000,000. If such amount exceeds US\$100,000,000, the referenced declaration must be submitted quarterly, in the month following the end of each quarter. Assets and rights that must be reported include the following: (i) bank deposits; (ii) loans; (iii) financing transactions; (iv) leases; (v) direct investments; (vi) portfolio investments, including Shares acquired under the Plan and the Program; (vii) financial derivatives investments; and (viii) other investments, such as real estate. Please note that foreign individuals holding Brazilian visas are considered Brazilian residents for purposes of this reporting requirement and must declare at least the assets held abroad that were acquired subsequent to the date of admittance as a resident of Brazil. Individuals holding assets and rights outside of Brazil valued at less than US\$1,000,000 are not required to submit a declaration.

Tax on Financial Transactions. Payments to foreign countries, repatriation of funds into Brazil, and the conversion associated with such fund transfers into local currency, may be subject to the Tax on Financial Transactions. It is your responsibility to comply with any applicable Tax on Financial Transactions arising from participation in the Plan and you should consult your personal tax advisor for additional details.

BULGARIA

Exchange Control Information. You will be required to file statistical forms with the Bulgarian National Bank annually regarding your receivables in bank accounts abroad as well as securities held abroad (e.g., Shares acquired under the Plan) if the total sum of all such receivables and securities equals or exceeds a certain threshold as of the previous calendar year-end. The reports are due by

March 31. You should contact your bank in Bulgaria for additional information regarding these requirements.

CANADA

TERMS AND CONDITIONS

Termination of Employment. This provision supplements Section IX(l) of the Agreement:

For purposes of the Performance Units and except as explicitly and minimally required under applicable legislation or expressly provided in this Agreement, (i) your employment will termination and (ii) your right, if any, to earn, seek damages in lieu of, vest in or otherwise benefit from or participate in any portion of the Performance Units or the Plan will be measured by and immediately terminate, as of the date you no longer actually providing services to the Company and/or the Employer, regardless of the reason for such termination and whether or not later found to be invalid or in breach of applicable laws in the jurisdiction where your employed or the terms of your employment agreement, if any (the “Termination Date”). Except as explicitly and minimally required under applicable legislation, the Termination Date will exclude and will not be extended by any period during which notice, pay in lieu of notice or related payments or damages are provided or required to be provided under applicable laws (including, but not limited to, statutory law, regulatory law and/or common or civil law). You will not earn or be entitled to any pro-rated vesting for that portion of time before the Termination Date, nor will you be entitled to any compensation for lost vesting, benefits, or other participation. Subject to applicable legislation, if the date you are no longer actually providing services cannot be reasonably determined under the terms of this Agreement or the Plan, the Company shall have the exclusive discretion to determine when you are no longer actually providing services for purposes of this Agreement (including whether you may still be considered to be providing services while on a leave of absence). Notwithstanding the foregoing, if applicable employment standards legislation explicitly requires continued vesting or other participation during a statutory notice period, your right to vest in the Performance Units or otherwise participate in or benefit from the Performance Units under the Plan, if any, will terminate effective as of the last day of your minimum statutory notice period, but you will not earn or be entitled to pro-rated vesting or other benefits or participation if the vesting date falls after the end of your statutory notice period, nor will you be entitled to any compensation for lost vesting;

Acknowledgement of Nature of Plan and Units. Section IX(c), IX(k) and IX(o)(A) of the Agreement are amended to read as follows:

(c) no Affiliate (including, but not limited to, the Employer) has any obligation to make any payment of any kind to you under this Agreement, except to the extent explicitly and minimally required under applicable legislation;

(k) in consideration of the grant of Performance Units hereunder, no claim or entitlement to compensation or damages arises from termination of Performance Units, and no claim

or entitlement to compensation or damages shall arise from forfeiture of the Performance Units resulting from (i) the termination of your employment by the Company or an Affiliate (regardless of the reason for such termination and whether or not later found to be invalid or in breach of employment laws in the jurisdiction where you are employed or the terms of your employment agreement, if any) and/or (ii) the application of the Clawback Policy as described in Section XXI below, except to the extent explicitly and minimally required under applicable legislation;

(o)(A) for employment law purposes outside the United States, the Performance Units, the Shares subject to the Performance Units, and the income from and value of same, are not part of normal or expected compensation or salary for any purpose, including but not limited to for purposes of calculating any severance, resignation, termination, redundancy, dismissal, end of service payments, bonuses, holiday pay, long-service awards, pension or retirement benefits or similar mandatory payments, except to the extent explicitly and minimally required under applicable legislation; and

Form of Settlement - Performance Units Payable Only in Shares. Notwithstanding any discretion in Section 9.5 of the Plan or the Program or anything to the contrary in the Agreement, the Award does not provide any right for you, as a resident of Canada, to receive a cash payment and shall be paid in Shares only.

The following provision will apply to you if you are a resident of Quebec:

French Language Documents. A French translation of this document and certain other documents related to this Award will be made available to you as soon as reasonably practicable. You understand that, from time to time, additional information related to the Award may be provided in English and such information may not be immediately available in French. However, upon request, the Company will provide a translation of such information into French as soon as reasonably practicable. Notwithstanding anything to the contrary in the Agreement, and unless you indicate otherwise, the French translation of this document and certain other documents related to the Award will govern your participation in the Plan.

Data Privacy Notice. This provision supplements Section XV of the Agreement:

You hereby authorize the Company and the Company's representative to discuss with and obtain all relevant information from all personnel (professional or not) involved in the administration of the Plan and the Program. You further authorize the Company, your Employer and Merrill Lynch Bank & Trust Co., FSB (or any other stock plan service provider) to disclose and discuss your participation in the Plan with their advisors. You also authorize the Company and your Employer to record such information and keep it in your file.

NOTIFICATIONS

Securities Law Information. You are permitted to sell Shares acquired through the Plan through the designated broker appointed under the Plan, if any, provided that the resale of such Shares takes

place outside of Canada through the facilities of a stock exchange on which the Shares are listed (e.g., the Nasdaq Global Select Market).

Foreign Asset/Account Reporting Information. Specified foreign property, including Shares, stock options and other rights to receive Shares (e.g., Performance Units) of a non-Canadian company held by a Canadian resident employee generally must be reported annually on a Form T1135 (Foreign Income Verification Statement) if the total cost of the employee's specified foreign property exceeds a certain threshold (currently C\$100,000) at any time during the year. Thus, such stock options and Performance Units must be reported – generally at nil cost – if the applicable cost threshold is exceeded because other specified foreign property is held by the employee. When Shares are acquired, their cost generally is the adjusted cost base (“ACB”) of the Shares. The ACB ordinarily would equal the fair market value of the Shares at the time of acquisition, but if the employee owns other shares of the same company, this ACB may have to be averaged with the ACB of the other shares.

CHINA

TERMS AND CONDITIONS

The following terms apply only to individuals who are subject to exchange control restrictions in the People's Republic of China (the “PRC”), as determined by the Company in its sole discretion:

Vesting of the Performance Units. Notwithstanding anything to the contrary in Article 7.1 of the Program, if your employment with the Company or an Affiliate terminates at any time during the Performance Period, you shall forfeit all Performance Units.

Sale Requirement. Notwithstanding anything to the contrary in the Agreement, due to exchange control laws in the PRC, you agree that the Company reserves the right to require the immediate sale of any Shares acquired upon settlement of the Performance Units. You understand and agree that any such immediate sale of Shares will occur as soon as is practical following settlement of the Performance Units. Alternatively, if the Shares are not immediately sold upon settlement of the Performance Units, the Company will require the sale of any Shares you may then hold within six (6) months (or such other period as may be required under applicable legal or exchange control requirements) following the termination of your employment with the Company, including its Affiliates.

You agree that the Company is authorized to instruct Merrill Lynch Bank & Trust Co., FSB or such other designated broker as may be selected by the Company to assist with the sale of the Shares on your behalf pursuant to this authorization, and you expressly authorize such broker to complete the sale of such Shares. You also agree to sign any agreements, forms and/or consents that may be reasonably requested by the Company (or the Company's designated broker) to effectuate the sale of the Shares (including, without limitation, as to the transfers of the proceeds and other exchange control matters noted below) and to otherwise cooperate with the Company with respect to such matters, provided that you shall not be permitted to exercise any influence

over how, when or whether the sales occur. Upon the sale of the Shares, you will receive the cash proceeds from the sale, less any applicable Tax Obligations, brokerage fees or commissions, in accordance with applicable exchange control laws and regulations.

You acknowledge that Merrill Lynch Bank & Trust Co., FSB or such other designated broker as may be selected by the Company is under no obligation to arrange for the sale of the Shares at any particular price. Due to fluctuations in the Share price and/or applicable exchange rates between the settlement date and (if later) the date on which the Shares are sold, the amount of proceeds ultimately distributed to you may be more or less than the market value of the Shares on the settlement date (which is the amount relevant to determining your liability for Tax Obligations). You understand and agree that the Company is not responsible for the amount of any loss that you may incur and that the Company assumes no liability for any fluctuations in the Share price and/or any applicable exchange rate.

Designated Broker Account. If Shares issued upon the settlement of the Performance Units are not immediately sold, you acknowledge that you are required to maintain the Shares in an account with Merrill Lynch Bank & Trust Co., FSB or such other designated broker as may be selected by the Company until the Shares are sold through such Company-designated broker.

Exchange Control Requirements. You understand and agree that, pursuant to local exchange control requirements, you will be required to repatriate the cash proceeds from the sale of the Shares issued upon settlement of the Performance Units and from the receipt of any Dividends or Dividend Equivalents to China. You further understand that, under applicable laws, such repatriation of your cash proceeds will need to be effectuated through a special exchange control account established by the Company or any Affiliate, including your Employer, and you hereby consent and agree that any proceeds may be transferred to such special account prior to being delivered to you. You also understand that the Company will deliver the proceeds to you as soon as possible, but that there may be delays in distributing the funds to you due to exchange control requirements in China. Proceeds may be paid to you in U.S. dollars or local currency at the Company's discretion. If the proceeds are paid to you in U.S. dollars, you will be required to set up a U.S. dollar bank account in China so that the proceeds may be deposited into this account. If the proceeds are paid to you in local currency, the Company is under no obligation to secure any particular currency conversion rate and the Company may face delays in converting the proceeds to local currency due to exchange control restrictions. You further agree to comply with any other requirements that may be imposed by the Company in the future in order to facilitate compliance with exchange control requirements in China.

COLOMBIA

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section IX of the Agreement:

You acknowledge that pursuant to Article 15 of Law 50/1990 (Article 128 of the Colombian Labor Code), the Plan, the Program and related benefits do not constitute a component of “salary” for any purpose. Therefore, they are considered to be of an extraordinary nature and will not be included and/or

considered for purposes of calculating any and all labor benefits, such as legal/fringe benefits, vacations, indemnities, payroll taxes, social insurance contributions and/or any other labor-related amounts, subject to the limitations provided in Law 1393/2010.

Mandate Letter. In accepting the Performance Units, you agree that – if requested by the Company or the Employer – you will execute a Mandate Letter or such other document (whether electronically or by such other method as requested by the Company or the Employer) that the Company determines is necessary or advisable in order that (i) a sufficient number of Shares to be allocated to you upon vesting can be sold on your behalf to cover Tax Obligations required to be withheld by the Employer and (ii) the proceeds from such sale can be wired directly from the Company to the Employer in Colombia for remittance to the tax authorities.

NOTIFICATIONS

Securities Law Information. The Shares are not and will not be registered with the Colombian registry of publicly traded securities (*Registro Nacional de Valores y Emisores*) and therefore the Shares may not be offered to the public in Colombia. Nothing in this document should be construed as the making of a public offer of securities in Colombia.

Exchange Control Information. Investment in assets located abroad (such as Shares acquired under the Plan and the Program) does not require prior approval from the Central Bank (Banco de la República). Nonetheless, such investments are subject to registration before the Central Bank as foreign investments held abroad, regardless of value.

Any payments for your investment originating in Colombia (and the liquidation of such investments) must be transferred through the Colombian foreign exchange market (*e.g.*, local banks), which includes the obligation to correctly complete and file the appropriate foreign exchange form (*declaración de cambio*).

Foreign Asset/Account Reporting Notice. An annual information return may need to be filed with the Colombian Tax Office detailing any assets held abroad (including Shares acquired under the Plan). If the individual value of any of these assets exceeds a certain threshold, each asset must be described (*e.g.*, its nature and its value) and the jurisdiction in which it is located must be disclosed. It is your responsibility to comply with this tax reporting requirement.

CROATIA

NOTIFICATIONS

Exchange Control Information. Croatian residents may be required to report any foreign investments (including Shares acquired under the Plan) to the Croatian National Bank for statistical purposes. You should be aware that exchange control regulations in Croatia are subject to frequent change and you are solely responsible for ensuring your continued compliance with current Croatian exchange control laws.

CZECH REPUBLIC

NOTIFICATIONS

Exchange Control Information. If you are a resident of the Czech Republic, you may be required to notify the Czech National Bank (“**CNB**”) of the acquisition of Shares under the Plan or maintenance of a foreign account if (i) you maintain foreign direct investments with a value of 2,500,000 Kč or more in the aggregate, (ii) you maintain a certain threshold of foreign financial assets, or (iii) you are specifically requested to do so by the CNB.

DENMARK

TERMS AND CONDITIONS

Danish Stock Option Act. In accepting the Performance Units, you acknowledge that you have received an Employer Statement translated into Danish, which is being provided to comply with the Danish Stock Option Act (the “Act”). To the extent more favorable to you and required to comply with the Act, as amended with effect from January 1, 2019. You also acknowledge any grant of Performance Units under the Plan made on or after January 1, 2019 is subject to the rules of the amended Act. Accordingly, you agree that the treatment of the Performance Units upon the termination of your employment is governed solely by Section I(i) of the Agreement and any corresponding provisions in the Plan. The relevant termination provisions are also detailed in the Employer Statement.

Please be aware that as set forth in Section 1 of the Act, the Act only applies to “employees” as that term is defined in Section 2 of the Act. If you are a member of the registered management of an Affiliate or Subsidiary in Denmark or otherwise do not satisfy the definition of employee, you will not be subject to the Act, and the Employer Statement will not apply to you.

NOTIFICATIONS

Foreign Asset/Account Reporting Information. The requirement to report certain information to the Danish Tax Administration via Form V or K was eliminated effective January 1, 2019. However, you still must report the foreign bank/brokerage accounts and their deposits, and Shares held in a foreign bank or brokerage account in your tax return under the section on foreign affairs and income.

EGYPT

NOTIFICATIONS

Exchange Control Information. If you transfer funds into Egypt in connection with the Performance Units, you are required to transfer the funds through a registered bank in Egypt.

FINLAND

NOTIFICATIONS

Foreign Asset/Account Reporting Information. There are no specific reporting requirements with respect to foreign assets/accounts. However, please note that you must check your pre-completed tax return to confirm that the ownership of Shares and other securities (foreign or domestic) are correctly reported. If you find any errors or omissions, you must make the necessary corrections electronically or by sending specific paper forms to the local tax authorities.

FRANCE

TERMS AND CONDITIONS

Language Consent. By accepting the Award, you confirm having read and understood the Plan and Agreement which were provided in the English language. You accept the terms of these documents accordingly.

Consentement Relatif à la Langue Utilisée. *En acceptant l'prix, vous confirmez avoir lu et compris le Plan et le Contrat, qui ont été communiqués en langue anglaise. Vous acceptez les termes de ces documents en connaissance de cause.*

NOTIFICATIONS

Foreign Asset/Account Reporting Information. French residents and non-residents must declare to the Customs Authorities the cash and securities they import or export without the use of a financial institution when the value of such cash or securities exceeds a certain threshold (currently €10,000). French residents also must report all foreign bank and brokerage accounts on an annual basis (including accounts opened or closed during the tax year) on Form N° 3916, together with the income tax return. Failure to comply could trigger significant penalties.

GERMANY

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If your acquisition of Shares under the Plan leads to a qualified participation at any point during the calendar year, you will need to report the acquisition when you file your tax return for the relevant year. A qualified participation is attained only in the unlikely event (i) you own at least 1% of the Company and the value of the Shares acquired exceeds a certain threshold (currently €150,000) or (ii) you hold Shares exceeding 10% of the Company's total Common Stock.

Exchange Control Information. Cross-border payments in excess of a certain threshold (currently €50,000) must be reported monthly to the German Federal Bank (*Bundesbank*). In case of payments in connection with securities (including proceeds realized upon the sale of Shares or the receipt of Dividends or Dividend Equivalents), and/or if the Company withholds Shares with a value in excess of the applicable threshold for any Tax Obligations, the report must be made monthly following the month in which the payment was received and must be filed electronically. The form of report (*Allgemeines Meldeportal Statistik*) can be accessed via the Bundesbank's website (www.bundesbank.de) and is

available in both German and English. In addition, you may be required to report the acquisition or sale of Shares to the *Bundesbank* if the value of the Shares acquired or sold exceeds the applicable threshold. You are responsible for satisfying any applicable reporting obligation.

GREECE

NOTIFICATIONS

Foreign Asset/Account Reporting Information. The reporting of foreign assets (including Shares and other investments) is your own obligation and takes place through your annual tax return.

HONG KONG

TERMS AND CONDITIONS

Form of Settlement - Performance Units Payable Only in Shares. Notwithstanding any discretion in Section 9.5 of the Plan or the Program or anything to the contrary in the Agreement, the Award does not provide any right for you, as a resident of Hong Kong, to receive a cash payment and shall be paid in Shares only.

Sale of Shares. Shares received at vesting are accepted as a personal investment. In the event that Shares are issued in respect of Performance Units within six (6) months of the Grant Date, you agree that you will not offer to the public or otherwise dispose of such Shares prior to the six (6)-month anniversary of the Grant Date.

NOTIFICATIONS

SECURITIES WARNING: *The contents of this document have not been reviewed by any regulatory authority in Hong Kong. You should exercise caution in relation to the offer. If you are in doubt about any of the contents of the Agreement, including this Appendix, or the Plan, you should obtain independent professional advice. The Performance Units and any Shares issued in respect of the Performance Units do not constitute a public offering of securities under Hong Kong law and are available only to members of the Board and Employees. The Agreement, including this Appendix, the Plan and other incidental communication materials have not been prepared in accordance with and are not intended to constitute a “prospectus” for a public offering of securities under the applicable securities legislation in Hong Kong. The Performance Units and any documentation related thereto are intended solely for the personal use of each member of the Board and/or Employee and may not be distributed to any other person.*

HUNGARY

There are no country-specific provisions.

ICELAND

NOTIFICATIONS

Exchange Control Information. Approval by the Central Bank of Iceland is no longer required to participate in the Plan and the Program, regardless of the value of the Shares acquired under the Plan and the Program. Despite the recent relaxation of the exchange control requirements, you should consult with your personal advisor to ensure compliance with applicable exchange control regulations in Iceland as such regulations are subject to frequent change. You are responsible for ensuring compliance with all exchange control laws in Iceland.

INDIA

NOTIFICATIONS

Exchange Control Information. You understand that you must repatriate any cash Dividends paid on Shares acquired under the Plan to India or any Dividend Equivalents paid in cash, as well as any proceeds from the sale of Shares acquired under the Plan within such time as may be required under applicable Indian exchange control laws, which may be amended from time to time, unless an exemption applies. You will receive a foreign inward remittance certificate (“FIRC”) from the bank where you deposit the foreign currency, and you must maintain the FIRC as proof of repatriation of funds in the event that the Reserve Bank of India or your Employer requests proof of repatriation. It is your responsibility to comply with these requirements. Neither the Company nor the Employer will be liable for any fines or penalties resulting from your failure to comply with any applicable laws. You may be required to provide information regarding funds received from participation in the Plan to the Company and/or the Employer to enable them to comply with their filing requirements under exchange control laws in India.

Foreign Asset/Account Reporting Information. You are required to declare foreign bank accounts and any foreign financial assets (including Shares held outside of India) in your annual tax return. It is your responsibility to comply with this reporting obligation, and you should consult your personal tax advisor in this regard.

IRELAND

TERMS AND CONDITIONS

Nature of Grant. This provision supplements Section IX of the Agreement:

In accepting the grant of Performance Units, you acknowledge that the benefits received under the Plan will not be taken into account for any redundancy or unfair dismissal claim.

ITALY

TERMS AND CONDITIONS

Nature of Grant. In accepting the grant of Performance Units, you acknowledge that (1) you have received a copy of the Plan, the Program, the Agreement and this Appendix; (2) you have reviewed the applicable documents in their entirety and fully understand the contents thereof; and (3) you accept all provisions of the Plan, the Program, the Agreement and this Appendix.

You further acknowledge that you have read and specifically and explicitly approve, without limitation, the following sections of the Agreement: Section III, Section IV, Section V, Section IX, Section IV, Section XVI, Section XX and the Data Privacy Notice for All European Economic Area (“EEA”) / European Union (“EU”) Jurisdictions, United Kingdom and Switzerland in this Appendix.

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Italian residents who, at any time during the fiscal year, hold foreign financial assets (including cash and Shares) which may generate income taxable in Italy are required to report these assets on their annual tax returns (UNICO Form, RW Schedule) for the year during which the assets are held, or on a special form if no tax return is due. These reporting obligations will also apply to Italian residents who are the beneficial owners of foreign financial assets under Italian money laundering provisions.

Foreign Financial Assets Tax. The fair market value of any Shares held outside of Italy is subject to a foreign assets tax at a flat rate. The market value is considered to be the value of the Shares on the Nasdaq Global Select Market on December 31 of the applicable year in which you held the Shares (or when the Shares are acquired during the course of the year, the tax is levied in proportion to the actual days of holding over the calendar year). No tax payment duties arise if the amount of the foreign financial assets tax calculated on all financial assets held abroad does not exceed a certain threshold. You should consult with your personal tax advisor about the foreign financial assets tax.

JAPAN

NOTIFICATIONS

Foreign Asset/Account Reporting Information. You will be required to report to the Japanese tax authorities details of any assets held outside of Japan as of December 31st (including any Shares acquired under the Plan and the Program) to the extent such assets have a total net fair market value exceeding a certain threshold (currently ¥50,000,000). Such report will be due by June 30 each year. You should consult with your personal tax advisor as to whether the reporting obligation applies to you and whether you will be required to include in the report details of any Shares or cash that you hold.

KOREA

NOTIFICATIONS

Exchange Control Information. Korean residents who sell Shares acquired under the Plan and/or receive cash Dividends on the Shares must file a report with a Korean foreign exchange bank if the

proceeds exceed a certain threshold (currently US\$5,000 per transaction) and are deposited into a non- Korean bank account. The report is not required if proceeds are deposited into a non-Korean brokerage account. It is your responsibility to ensure compliance with any applicable exchange control reporting obligations.

Foreign Asset/Account Reporting Information. You are required to declare all foreign financial accounts (*e.g.* non-Korean bank accounts, brokerage accounts holding Shares, etc.) to the Korean tax authority and file a report regarding such accounts if the monthly balance of such accounts exceeds a certain threshold. It is your responsibility to comply with this reporting obligation and you should consult your personal tax advisor to ensure compliance with this requirement.

LITHUANIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. If you (i) hold certain job positions established by the law or (ii) donate to political parties or political campaigners, you must file an Annual Asset Return of the Individual (Family) in Form No. FR0001 with respect to assets held outside of Lithuania (*e.g.*, Shares). If you open an account in a foreign financial institution and annual turnover in the account exceeds a certain threshold (currently €15,000), you must file a foreign account report.

MEXICO

TERMS AND CONDITIONS

Acknowledgement of the Grant. In accepting the Award granted hereunder, you acknowledge that you have received a copy of the Plan and the Program, have reviewed the Plan and the Program and the Agreement, including this Appendix, in their entirety and fully understand and accept all provisions of the Plan, the Program and the Agreement, including this Appendix. You further acknowledge that you have read and specifically and expressly approve the terms and conditions of Section IX of the Agreement, in which the following is clearly described and established:

- (1) Your participation in the Plan and the Program do not constitute an acquired right.
- (2) The Plan and your participation in the Plan and the Program are offered by Amgen Inc. on a wholly discretionary basis.
- (3) Your participation in the Plan and the Program is voluntary.
- (4) Amgen Inc. and its Affiliates are not responsible for any decrease in the value of any Shares issued with respect to the Award.

Labor Law Acknowledgement and Policy Statement. In accepting any Award granted hereunder, you expressly recognize that Amgen Inc., with registered offices at One Amgen Center Drive, Thousand Oaks, California 91320, U.S.A., is solely responsible for the administration of the Plan and

that your participation in the Plan and acquisition of Shares do not constitute an employment relationship between you and Amgen Inc. since you are participating in the Plan on a wholly commercial basis and your sole employer is Amgen Mexico S.A. de C.V. ("Amgen-Mexico"). Based on the foregoing, you expressly recognize that the Plan and the Program and the benefits that you may derive from participation in the Plan and the Program do not establish any rights between you and your Employer, Amgen-Mexico, and do not form part of the employment conditions and/or benefits provided by Amgen-Mexico and any modification of the Plan or its termination shall not constitute a change or impairment of the terms and conditions of your employment.

You further understand that your participation in the Plan and the Program is as a result of a unilateral and discretionary decision of Amgen Inc.; therefore, Amgen Inc. reserves the absolute right to amend and/or discontinue your participation in the Plan at any time without any liability to you.

Finally, you hereby declare that you do not reserve to yourself any action or right to bring any claim against Amgen Inc. for any compensation or damages regarding any provision of the Plan or the benefits derived under the Plan, and you therefore grant a full and broad release to Amgen Inc., its Affiliates, stockholders, officers, agents or legal representatives with respect to any claim that may arise.

Spanish Translation

Reconocimiento del Otorgamiento. Al aceptar cualquier Otorgamiento de Acciones bajo el presente documento, usted reconoce que ha recibido una copia del Plan y del Programa, que ha revisado el Plan y el Programa, así como también el Apéndice en su totalidad, además que comprende y está de acuerdo con todas las disposiciones tanto del Plan, del Programa y del Otorgamiento, incluyendo este Apéndice. Asimismo, usted reconoce que ha leído y manifiesta específicamente y expresamente la conformidad con los términos y condiciones establecidos en la Sección IX del Acuerdo del Otorgamiento, en los que se establece y describe claramente que:

- (1) Su participación en el Plan y en el Programa de ninguna manera constituye un derecho adquirido.
- (2) Su participación en Plan y en el Programa son ofrecidos por Amgen Inc. de forma completamente discrecional.
- (3) Su participación en el Plan y en el Programa es voluntaria.
- (4) Amgen Inc. y sus Afiliados no son responsables de ninguna disminución en el valor de las Acciones Comunes emitidas mediante el Plan.

Reconocimiento de la Ley Laboral y Declaración de Política. Al aceptar cualquier Otorgamiento bajo el presente, usted reconoce expresamente que Amgen Inc., con oficinas registradas localizadas en One Amgen Center Drive, Thousand Oaks, California 91320, U.S.A., es la única responsable de la administración del Plan y que su participación en el mismo y la adquisición de Acciones Comunes no constituyen de ninguna manera una relación laboral entre usted y Amgen Inc., debido a que su

participación en el Plan es únicamente una relación comercial y que su único empleador es Amgen Mexico S.A. de C.V. (“Amgen-Mexico”). Derivado de lo anterior, usted reconoce expresamente que el Plan y el Programa y los beneficios a su favor que pudieran derivar de la participación en el mismo, no establecen ningún derecho entre usted y su empleador, Amgen – México, y no forman parte de las condiciones laborales y/o los beneficios otorgados por Amgen – México, y cualquier modificación del Plan o la terminación del mismo no constituirá un cambio o desmejora de los términos y condiciones de su trabajo.

Asimismo, usted entiende que su participación en el Plan y en el Programa es resultado de la decisión unilateral y discrecional de Amgen Inc., por lo tanto, Amgen Inc. se reserva el derecho absoluto de modificar y/o discontinuar su participación en el Plan en cualquier momento y sin ninguna responsabilidad para usted.

Finalmente, usted manifiesta que no se reserva ninguna acción o derecho que origine una demanda en contra de Amgen Inc., por cualquier compensación o daños y perjuicios, en relación con cualquier disposición del Plan o de los beneficios derivados del mismo, y en consecuencia usted exime amplia y completamente a Amgen Inc. de toda responsabilidad, como así también a sus Afiliadas, accionistas, directores, agentes o representantes legales con respecto a cualquier demanda que pudiera surgir.

NOTIFICATIONS

Securities Law Information. The Performance Units and the Shares offered under the Plan have not been registered with the National Register of Securities maintained by the Mexican National Banking and Securities Commission and cannot be offered or sold publicly in Mexico. In addition, the Plan, the Agreement and any other document relating to the Performance Units may not be publicly distributed in Mexico. These materials are addressed to you only because of your existing relationship with the Company and your Employer and these materials should not be reproduced or copied in any form. The offer contained in these materials does not constitute a public offering of securities but rather constitutes a private placement of securities addressed specifically to individuals who are present employees of Amgen-Mexico made in accordance with the provisions of the Mexican Securities Market Law, and any rights under such offering shall not be assigned or transferred.

NETHERLANDS

NOTIFICATIONS

Securities Law Information.

**Attention! This investment falls outside AFM supervision.
No prospectus required for this activity.**



NORWAY

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Norwegian residents may be subject to foreign asset reporting as part of their ordinary tax return. Norwegian banks, financial institutions, limited companies etc. must report certain information to the Tax Administration. Such information may then be pre-completed in a Norwegian resident's tax return. However, if the resident has traded, or is the owner of, financial instruments (e.g., Shares) not pre-completed in the tax return, the Norwegian resident must enter this information in Form RF-1159, which is an appendix to the tax return.

Exchange Control Information. In general, Norwegian residents should not be subject to any foreign exchange requirements in connection with their acquisition or sale of Shares under the Plan, except normal reporting requirements to the Norwegian Currency Registry. If any transfer of funds into or out of Norway is made through a Norwegian bank, the bank will make the registration.

POLAND

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Polish residents holding foreign securities (including Shares) and maintaining accounts abroad must file reports with the National Bank of Poland if the aggregate value of cash and securities held in such foreign accounts exceeds a certain threshold (currently PLN 7 million). If required, the reports are due on a quarterly basis by the 20th day following the end of each quarter and must be filed on special forms available on the website of the National Bank of Poland.

Exchange Control Information. In addition, Polish residents are required to transfer funds through a bank account in Poland if the transferred amount in any single transaction exceeds a specified threshold (currently €15,000 (or PLN 15,000 if such transfer of funds is associated with the business activity of a consultant)). You must store all documents connected with any foreign exchange transactions you engage in for a period of five (5) years from the end of the year when such transactions were made. Penalties may apply for failure to comply with exchange control requirements.

PORTUGAL

TERMS AND CONDITIONS

Consent to Receive Information in English. You hereby expressly declare that you have full knowledge of the English language and have read, understood and fully accepted and agreed with the terms and conditions established in the Plan, the Program and Agreement.

Conhecimento da Língua. *Por meio do presente, eu declaro expressamente que tem pleno conhecimento da língua inglesa e que li, compreendi e livremente aceitei e concordei com os termos e condições estabelecidas no Plano, no Programa e no Acordo.*

PUERTO RICO

There are no country-specific provisions.

ROMANIA

NOTIFICATIONS

Exchange Control Information. Certain transfers of funds may need to be reported to the National Office for Prevention and Control of Money Laundering on specific forms by the relevant bank or financial institution. If you deposit proceeds from the sale of Shares or the receipt of Dividends or Dividend Equivalents in a bank account in Romania, you may be required to provide the Romanian bank assisting with the transaction with appropriate documentation explaining the source of the income. You should consult with a legal advisor to determine whether you will be required to submit such documentation to the Romanian bank.

RUSSIA

NOTIFICATIONS

Foreign Account Report Information. Russian residents maintaining accounts abroad must disclose such accounts with the Russian tax authorities, as follows: (i) file notifications upon opening (or closing or changing bank details) within 30 days, and (ii) file reports on the movement of cash or financial assets on such accounts by June 1 of the following year. Such notifications and reports must be filed on

special forms available on the website of the Russian tax authorities. Russian residents who are absent from Russia for more than 183 days in a year may be relieved from this reporting.

TERMS AND CONDITIONS

Securities Law Information. The sale of Shares acquired under the Plan is not permitted within Russia. The Shares are currently traded on the Nasdaq Global Select Market in the U.S. under the ticker symbol “AMGN,” therefore, the Shares may be sold on this exchange.

You understand that the exchange control rules and regulations in Russia and the legal restrictions impacting your participation in the Plan are subject to frequent change. You should consult with your personal legal advisor to determine the applicability of any requirements or restrictions applicable to any Shares or cash received in connection with the Plan.

SINGAPORE

TERMS AND CONDITIONS

Restriction on Sale and Transferability. You hereby agree that any Shares acquired pursuant to the Performance Units will not be offered for sale in Singapore prior to the six (6)-month anniversary of the Grant Date, unless such sale or offer is made pursuant to one or more exemptions under Part XIII Division 1 Subdivision (4) (other than section 280) of the Securities and Futures Act (Chap. 289, 2006 Ed.) (“SFA”), or pursuant to, and in accordance with the conditions of, any other applicable provisions of the SFA.

NOTIFICATIONS

Securities Law Information. The grant of the Performance Units is being made pursuant to the “Qualifying Person” exemption under section 273(1)(f) of the SFA, on which basis it is exempt from the prospectus and registration requirements under the SFA, and is not made with a view to the Performance Units being subsequently offered for sale to any other party. The Plan has not been, and will not be, lodged or registered as a prospectus with the Monetary Authority of Singapore.

Director Notification Requirement. Directors (including alternate, substitute, associate and shadow directors) of a Singapore Affiliate are subject to certain notification requirements under the Singapore Companies Act, regardless of whether they are resident or employed in Singapore. Directors of a Singapore Affiliate must notify the Singapore Affiliate in writing of an interest (*e.g.*, Performance Units, Shares, etc.) in the Company or any related company within two (2) business days of (i) its acquisition or disposal, (ii) any change in a previously disclosed interest (*e.g.*, when the Shares are sold), or (iii) becoming a director.

SLOVAK REPUBLIC

There are no country-specific provisions.

SLOVENIA

NOTIFICATIONS

Foreign Asset/Account Reporting Information. Slovenian residents may be required to report the opening of bank and/or brokerage accounts to tax authorities within eight (8) days of opening such account. You should consult with your personal tax advisor to determine whether this requirement will be applicable to any accounts opened in connection with your participation in the Plan (*e.g.*, your brokerage account with the Company's designated broker).

SPAIN

TERMS AND CONDITIONS

Labor Law Acknowledgement. The following provision supplements Section IX of the Agreement:

By accepting the Award granted hereunder, you consent to participation in the Plan and the Program and acknowledge that you have received a copy of the Plan and the Program.

You understand that the Company has unilaterally, gratuitously and in its sole discretion decided to grant the Award under the Plan and the Program to individuals who may be members of the Board or Employees of the Company or its Affiliates throughout the world. The decision is a limited decision that is entered into upon the express assumption and condition that (i) the Awards granted will not economically or otherwise bind the Company or any of its Affiliates on an ongoing basis, other than as expressly set forth in the applicable Agreement, including this Appendix; (ii) the Awards granted hereunder and any Shares acquired under the Plan shall not be considered a mandatory benefit, salary for any purposes (including severance compensation) or any other right whatsoever; and (iii) any unvested Performance Units will be forfeited immediately upon termination of your employment for any reason, as detailed below. Further, you understand and freely accept that there is no guarantee that any benefit whatsoever shall arise from any gratuitous and discretionary grant of the Award since the future value of the Award and any Shares that may be issued in respect of such Award is unknown and unpredictable. In addition, you understand that the Award granted hereunder would not be made but for the assumptions and conditions referred to above; thus, you understand, acknowledge and freely accept that, should any or all of the assumptions be mistaken or should any of the conditions not be met for any reason, then the grant of the Award or right to the Award shall be null and void.

Further, the vesting of the Performance Units is expressly conditioned your continued and active rendering of service, such that if your employment terminates for any reason whatsoever, the Performance Units may cease vesting immediately, in whole or in part, effective on the date of your termination of employment (unless otherwise specifically provided in Section I of the Agreement). This will be the case, for example, even if (1) you are considered to be unfairly dismissed without good cause (*i.e.*, subject to a “despido improcedente”); (2) you are dismissed for disciplinary or objective reasons or due to a collective dismissal; (3) you terminate service due to a change of work location, duties or any other employment or contractual condition; (4) you terminate service due to a unilateral breach of contract by the Company or an Affiliate; or (5) your employment terminates for any other

reason whatsoever. Consequently, upon termination of your employment for any of the above reasons, you may automatically lose any rights to Performance Units that were not vested on the date of your termination of employment, as described in the Plan and the Agreement.

You acknowledge that you have read and specifically accept the conditions referred to in Section I of the Agreement.

NOTIFICATIONS

Securities Law Information. No “offer of securities to the public,” as defined under Spanish law, has taken place or will take place in the Spanish territory. The Agreement (including this Appendix) has not been nor will it be registered with the *Comisión Nacional del Mercado de Valores*, and does not constitute a public offering prospectus.

Exchange Control Information. You will be required to declare electronically to the Bank of Spain any security accounts (including brokerage accounts held abroad), as well as the securities (including Shares acquired under the Plan) held in such accounts if the value of the transactions for all such accounts during the prior year or the balances of such accounts as of December 31 of the prior year exceeds a certain threshold (currently, €1,000,000). Different thresholds and deadlines to file this declaration apply. However, if neither such transactions during the immediately preceding year nor the balances / positions as of December 31 exceed the applicable threshold, no such declaration must be filed unless expressly required by the Bank of Spain. If any of such thresholds were exceeded during the current year, you may be required to file the relevant declaration corresponding to the prior year, however, a summarized form of declaration may be available.

Foreign Asset/Account Reporting Information. To the extent that you hold Shares and/or have bank accounts outside of Spain with a value in excess of €50,000 (for each type of asset) as of December 31 each year, you will be required to report information on such assets in your tax return (tax form 720) for such year. After such Shares and/or accounts are initially reported, the reporting obligation will apply for subsequent years only if the value of any previously-reported Shares or accounts increases by more than €20,000 (for each type of asset) as of each subsequent December 31, or if you sell or otherwise dispose of previously-reported Shares or accounts. If the value of such Shares and/or accounts as of December 31 does not exceed €50,000, a summarized form of declaration may be presented.

SWEDEN

TERMS AND CONDITIONS

Authorization to Withhold. This provision supplements Section III of the Agreement:

Without limiting the Company’s and the Employer’s authority to satisfy their withholding obligations for Tax Obligations as set forth in the Agreement, in accepting the Performance Units, you authorize the Company to withhold Shares or to sell Shares otherwise issuable to you upon vesting or settlement to satisfy Tax Obligations, regardless of whether the Company and/or Employer have an obligation to

withhold such Tax Obligations, provided that such withholding would not, in the Company's determination, result in adverse accounting consequences to the Company.

SWITZERLAND

NOTIFICATIONS

Securities Law Information. Neither this document nor any other materials relating to the Performance Units (i) constitutes a prospectus according to articles 35 et seq. of the Swiss Federal Act on Financial Services ("FinSA"), (ii) may be publicly distributed or otherwise made publicly available to any person other than an employee of the Company or one of its Subsidiaries in Switzerland or (iii) has been or will be filed with, approved or supervised by any Swiss reviewing body according to article 51 of FinSA or any Swiss regulatory authority, including the Swiss Financial Market Supervisory Authority.

TAIWAN

NOTIFICATIONS

Securities Law Information. Participation in the Plan is available only for select Board members and Employees of the Company and its Affiliates. The grant of Performance Units and participation in the Plan is not a public offer of securities by a Taiwanese company.

Exchange Control Information. You may acquire and remit foreign currency (including proceeds from the sale of Shares or the receipt of Dividends or Dividend Equivalents) up to US\$10,000,000 per year without justification. If the transaction amount is TWD500,000 or more in a single transaction, you must submit a Foreign Exchange Transaction Form. If the transaction amount is US\$500,000 or more in a single transaction, you must also provide supporting documentation to the satisfaction of the remitting bank.

THAILAND

NOTIFICATIONS

Exchange Control Information. If proceeds from the sale of Shares or the receipt of any Dividends or Dividend Equivalents exceed a certain threshold (currently US\$1,000,000), you must (i) immediately repatriate such funds to Thailand and (ii) report the inward remittance to the Bank of Thailand on a Foreign Exchange Transaction Form. In addition, within three hundred and sixty (360) days of repatriation, you must either convert any funds repatriated to Thailand to Thai Baht or deposit the funds in a foreign exchange account with a Thai commercial bank. Any such commercial bank must be duly authorized by the Bank of Thailand to engage in the purchase, exchange and withdrawal of foreign currency.

TÜRKİYE

NOTIFICATIONS

Securities Law Information. The sale of Shares acquired under the Plan is not permitted within Türkiye. The sale of Shares acquired under the Plan must occur outside of Türkiye. The Shares are currently traded on the Nasdaq Global Select Market in the U.S. under the ticker symbol “AMGN” and Shares may be sold on this exchange.

Exchange Control Information. You may be required to engage a Turkish financial intermediary to assist with the sale of Shares acquired under the Plan. To the extent a Turkish financial intermediary is required in connection with the sale of any Shares acquired under the Plan, you are solely responsible for engaging such Turkish financial intermediary. You should consult your personal legal advisor prior to the sale of Shares to ensure compliance with the current requirements.

UNITED ARAB EMIRATES

NOTIFICATIONS

Securities Law Information. Performance Units under the Plan are available only to select Board members and Employees of the Company and its Affiliates and are for the purpose of providing equity incentives. The Plan, the Program and the Agreement are intended for distribution only to such Board members and Employees and must not be delivered to, or relied on by, any other person. You should conduct your own due diligence on the Performance Units offered pursuant to this Agreement. If you do not understand the contents of the Plan and/or the Agreement, you should consult an authorized financial adviser. The Emirates Securities and Commodities Authority and the Dubai Financial Services Authority have no responsibility for reviewing or verifying any documents in connection with the Plan. Further, the Ministry of the Economy and the Dubai Department of Economic Development have not approved the Plan or the Agreement nor taken steps to verify the information set out therein and have no responsibility for such documents.

UNITED KINGDOM

TERMS AND CONDITIONS

Tax Withholding. This provision supplements Section V of the Agreement:

Without limitation to Section V of the Agreement, you agree that you are liable for all Tax Obligations and hereby covenant to pay all such Tax Obligations as and when requested by the Company or your Employer or by HM Revenue and Customs (“HMRC”) (or any other tax authority or any other relevant authority). You also agree to indemnify and keep indemnified the Company and your Employer against any taxes that they are required to pay or withhold or have paid or will pay to HMRC (or any other tax authority or any other relevant authority) on your behalf.

Notwithstanding the foregoing, if you are an executive officer or director (as within the meaning of Section 13(k) of the Exchange Act, as amended, from time to time), you understand that you may not be able to indemnify the Company or your Employer for the amount of income tax not collected from or paid by you, as it may be considered a loan. In the event that you are an executive officer or director and income tax is not collected from you within ninety (90) days after the end of the tax year in which the Taxable Event occurs, the amount of any uncollected income tax may constitute an additional benefit to you on which additional income tax and national insurance contributions (“NICs”) may be payable. You acknowledge that you are responsible for reporting and paying any income tax due on this additional benefit directly to HMRC under the self-assessment regime and for paying your Employer for the amount of any NICs due on this additional benefit, which the Company or your Employer may obtain from you by any of the means set forth in Section V of the Agreement.

If the maximum applicable withholding rate is used, any over-withheld amount may be credited to you by the Company or your Employer (with no entitlement to the Common Stock equivalent) or if not so credited, you may seek a refund from the local tax authorities.

Joint Election. If you are a resident of the United Kingdom between the Grant Date and the vesting of the Performance Units, as a condition of the Award, you agree to accept any liability for secondary Class 1 National Insurance Contributions (the “Employer NICs”) which may be payable by the Company or your Employer with respect to the earning and/or payment of the Performance Units and issuance of Shares in respect of the Performance Units, the assignment or release of the Performance Units for consideration or the receipt of any other benefit in connection with the Performance Units.

Without limitation to the foregoing, you agree to make an election (the “Election”), in the form specified and/or approved for such election by HMRC, that the liability for your Employer NICs payments on any such gains shall be transferred to you to the fullest extent permitted by law. You further agree to execute such other elections as may be required between you and any successor to the Company and/or your Employer. You hereby authorize the Company and your Employer to withhold such Employer NICs by any of the means set forth in Section V of the Agreement.

Failure by you to enter into an Election, withdrawal of approval of the Election by HMRC or a joint revocation of the Election by you and the Company or your Employer, as applicable, shall be grounds for the forfeiture and cancellation of the Performance Units, without any liability to the Company or your Employer.

UNITED STATES

TERMS AND CONDITIONS

Nature of Grant. The following provision replaces Section IX(l) of the Award Agreement:

(l) in the event of termination of your employment (whether or not in breach of local labor laws), your right to receive Performance Units and receive Shares under the Plan and the Program, if any, will terminate effective as of the date that you are no longer actively employed; *provided, however*, that such right will be extended by any notice period mandated by law (e.g., the Worker Adjustment and

Retraining Notification Act (“WARN Act”) notice period or similar periods pursuant to local law) and any paid administrative leave (as applicable), unless the Company shall provide you with written notice otherwise before the commencement of such notice period or leave. In such event, payment of the Performance Units shall be made in accordance with Section IV; *provided, further, however*, that notwithstanding the effect of any such extension, subject to Section 4.2 of the Program, in no event will the Performance Units be paid later than the 90th day following the last day of the Performance Period.

CERTAIN CONFIDENTIAL INFORMATION CONTAINED IN THIS DOCUMENT, MARKED BY [*], HAS BEEN OMITTED BECAUSE IT IS BOTH (I) NOT MATERIAL AND (II) IS THE TYPE THAT THE REGISTRANT TREATS AS PRIVATE OR CONFIDENTIAL.

October 1, 2025

Amgen Inc.
One Amgen Center Drive
Thousand Oaks, California 91320-1799
United States
Attention: Corporate Secretary

Re: Waiver of Certain Rights Related to [*]-Targeting Molecules Under the Collaboration Agreement

Dear Sir or Madam:

Reference is made to that certain Collaboration Agreement, dated as of October 31, 2019 (as amended on April 20, 2022 and February 26, 2023, the "Collaboration Agreement"), by and among Amgen Inc. ("Amgen"), BeOne Medicines I GmbH (f/k/a BeiGene Switzerland GmbH) ("BeOne") and, solely with respect to Section 13.6, BeOne Medicines Ltd. (f/k/a BeiGene, Ltd.). Capitalized terms used but not defined herein shall have the meanings ascribed to them in the Collaboration Agreement.

Amgen has notified BeOne that Amgen is undergoing discussions with a potential counterparty ("Counterparty") regarding a [*] combination study with Amgen's IMDELLTRA[®] and Counterparty's [*] (the "Counterparty [*] Product" and the "Study").

In accordance with Section 15.16 (Waivers and Modifications) of the Collaboration Agreement, this letter serves to document BeOne's and Amgen's agreement that, with respect to the Counterparty [*] Product, the Distracting Program Restrictions under Article IX of the Collaboration Agreement are hereby waived by the Parties in relation to the Study.

Please acknowledge your agreement and consent to the foregoing by counter-signing this letter agreement in the space provided below and returning a copy to us.

[Signature Page Follows]

Yours sincerely,
BEONE MEDICINES I GMBH

/s/ Michael Schoen
By: Michael Schoen
Title: Managing Director

BEONE MEDICINES LTD.

/s/ Chan Lee
By: Chan Lee
Title: SVP, General Counsel and Corporate Secretary

Acknowledged and agreed:
AMGEN INC.

/s/ Rachna Khosla
By: Rachna Khosla
Title: SVP, Business Development

BeOne Medicines I GmbH
Aeschengraben 27
4051 Basel, Switzerland
BeOneMedicines.com

CERTAIN CONFIDENTIAL INFORMATION CONTAINED IN THIS DOCUMENT, MARKED BY [*], HAS BEEN OMITTED BECAUSE IT IS BOTH (I) NOT MATERIAL AND (II) IS THE TYPE THAT THE REGISTRANT TREATS AS PRIVATE OR CONFIDENTIAL.

THIRD AMENDMENT TO COLLABORATION AGREEMENT

This Third Amendment to the Collaboration Agreement (“Amendment”) is entered into as of October 31, 2025 (the “Third Amendment Effective Date”) by and among Amgen Inc., a Delaware corporation having its principal place of business at One Amgen Center Drive, Thousand Oaks, California 91320-1799 (“Amgen”), BeOne Medicines I GmbH (f/k/a BeiGene Switzerland GmbH), a Swiss corporation with a principal place of business at Aeschengraben 27, 4051 Basel, Switzerland (“BeOne”), and BeOne Medicines Ltd. (f/k/a BeiGene, Ltd.), a Swiss corporation with a principal place of business at Aeschengraben 27, 4051 Basel, Switzerland (“BeOne Parent”). BeOne and Amgen are sometimes referred to herein individually as a “Party” and collectively as the “Parties.” This Amendment amends that certain Collaboration Agreement, entered into as of October 31, 2019 (as amended from time to time, the “Agreement”), by and between Amgen and BeOne and, solely with respect to Section 13.6 thereof, BeOne Parent. Capitalized terms used but not defined herein have the meanings given to them in the Agreement.

RECITALS

WHEREAS, pursuant to the Agreement, Amgen and BeOne collaborate on the commercialization of certain Products (as defined in the Agreement) in the Collaboration Territory (as defined in the Agreement) and the global development funding and clinical development and commercialization of certain clinical-stage Pipeline Products in the Collaboration Territory; and

WHEREAS, the Parties desire to enter into this Amendment, upon the terms and conditions set forth herein.

NOW, THEREFORE, in consideration of the premises and the mutual covenants and agreements contained herein, the Parties, intending to be legally bound hereby, do agree as follows:

AGREEMENT

1. Amendments to Party Names.

- a. All references in the Agreement to “BeiGene Switzerland GmbH” are hereby amended to “BeOne Medicines I GmbH.”
- b. All references in the Agreement to “BeiGene, Ltd.” are hereby amended to “BeOne Medicines Ltd.”
- c. All references in the Agreement to “BeiGene” are hereby amended to “BeOne.”

d. All references in the Agreement to “BeiGene Parent” are hereby amended to “BeOne Parent.”

2. Amendment to Select Definitions.

a. The defined term “Bo Ao Product” is hereby amended and restated in its entirety as follows:

“Bo Ao Product” means any Product which is approved by the JSC for importation into Hainan Bo Ao prior to Regulatory Approval of such Product.”

b. The defined term “Early Access Product” is hereby amended and restated in its entirety as follows:

“Early Access Product” means each of a Bo Ao Product or a Tianjin Product.”

c. The defined term “Tianjin Product” is hereby amended and restated in its entirety as follows:

“Tianjin Product” means any Product which is approved by the JSC for importation into Tianjin prior to Regulatory Approval of such Product.”

d. The defined term “Tianjin Support Costs” is hereby amended and restated in its entirety as follows:

“Tianjin Support Costs” means all actual and, if reasonably practicable, [*].”

3. The first paragraph of Section 7.2.8 of the Agreement is hereby amended and restated in its entirety as follows:

“Notwithstanding anything to the contrary in this Agreement, the Parties desire to initiate the Profit-sharing arrangement set forth in Section 7.2 for any Bo Ao Product, Tianjin Product or Early Access Product prior to applicable Initiation Date, subject to the following terms and conditions:”

4. Supply Price Schedule. The Supply Price Schedule is hereby amended to include the following as a new row:

[*]	[*]%
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5. Miscellaneous.

a. Except as specifically amended above, the Agreement shall continue to be in full force and effect.

b. This Amendment and its effect are subject to and shall be construed and enforced in accordance with the laws of the State of New York, U.S.A.

c. This Amendment may be executed in counterparts with the same effect as if both Parties had signed the same document. All such counterparts will be deemed an original, will be construed together and will constitute one and the same instrument. Signature pages of this Amendment may be exchanged by facsimile or other electronic means without affecting the validity thereof.

IN WITNESS WHEREOF, the Parties have executed this Amendment as of the Third Amendment Effective Date.

AMGEN INC.

/s/ Rachna Khosla

Name: Rachna Khosla

Title: Senior Vice President, Business Development

BEONE MEDICINES I GMBH

/s/ Michael Schoen

Name: Michael Schoen

Title: Managing Director

BEONE MEDICINES LTD.

/s/ Chan Lee

Name: Chan Lee

Title: Senior Vice President, General Counsel & Corporate Secretary

CERTAIN CONFIDENTIAL INFORMATION CONTAINED IN THIS DOCUMENT, MARKED BY [*], HAS BEEN OMITTED BECAUSE IT IS BOTH (I) NOT MATERIAL AND (II) IS THE TYPE THAT THE REGISTRANT TREATS AS PRIVATE OR CONFIDENTIAL.

FOURTH AMENDMENT TO COLLABORATION AGREEMENT

This Fourth Amendment to the Collaboration Agreement (“Amendment”) is entered into as of November 11, 2025 (the “Fourth Amendment Effective Date”) by and among Amgen Inc., a Delaware corporation having its principal place of business at One Amgen Center Drive, Thousand Oaks, California 91320-1799 (“Amgen”), BeOne Medicines I GmbH (f/k/a BeiGene Switzerland GmbH), a Swiss corporation with a principal place of business at Aeschengraben 27, 4051 Basel, Switzerland (“BeOne”), and BeOne Medicines Ltd. (f/k/a BeiGene, Ltd.), a Swiss corporation with a principal place of business at Aeschengraben 27, 4051 Basel, Switzerland (“BeOne Parent”). BeOne and Amgen are sometimes referred to herein individually as a “Party” and collectively as the “Parties.” This Amendment amends that certain Collaboration Agreement, entered into as of October 31, 2019 (as amended from time to time, the “Agreement”), by and between Amgen and BeOne and, solely with respect to Section 13.6 thereof, BeOne Parent. Capitalized terms used but not defined herein have the meanings given to them in the Agreement.

RECITALS

WHEREAS, pursuant to the Agreement, Amgen and BeOne collaborate on the commercialization of certain Products (as defined in the Agreement) in the Collaboration Territory (as defined in the Agreement) and the global development funding and clinical development and commercialization of certain clinical-stage Pipeline Products in the Collaboration Territory; and

WHEREAS, the Parties desire to enter into this Amendment, upon the terms and conditions set forth herein.

NOW, THEREFORE, in consideration of the premises and the mutual covenants and agreements contained herein, the Parties, intending to be legally bound hereby, do agree as follows:

AGREEMENT

1. Amendment to Section 5.1.5(a) (Retained In-Line Product) of the Agreement. Section 5.1.5(a) of the Agreement is hereby amended and restated in its entirety as follows:

“BeOne shall retain [*] in the Collaboration Territory for so long as such In-Line Product is sold in the Collaboration Territory (each, a “Retained In-Line Product”).”

2. Amendment to Section 7.2.1 (BeOne Costs). Section 7.2.1 of the Agreement is hereby amended and restated in its entirety as follows:

““BeOne Costs. Within [*] after the end of each calendar quarter, BeOne will provide to Amgen a final report of its Commercialization and Related Costs, on a Product-by-

Product basis, incurred by BeOne or its Affiliates in accordance with this Agreement (collectively, “BeOne Costs”) in such quarter. BeOne will initially incur the portion of the Commercialization and Related Costs attributed to its activities hereunder. In addition to the annual Commercialization Budget approved hereunder, prior to the end of each calendar year, BeOne will provide Amgen with a nonbinding estimate of its Commercialization and Related Costs for the [*] period [*] following the year covered by such approved budgets; *provided* that the Parties will review and discuss such estimated Costs at the JSC. Notwithstanding the foregoing, BeOne’s shared portion of the Commercialization and Related Costs incurred in performing any Clinical Studies conducted after Regulatory Approval for In-Line Products in the Collaboration Territory shall be subject to an annual maximum of [*] during each [*] period following the Effective Date and until January 1, 2026 subject to an aggregate maximum of [*]. Within [*] after the end of each calendar quarter, BeOne will provide Amgen with a report of BeOne’s product-level profit & loss statements for such calendar quarter, which report will contain a detailed and itemized calculation of Net Revenues for each Product during such calendar quarter. Additionally, within [*] after the end of each calendar quarter, BeOne will provide Amgen with a report of any Recoveries for such calendar quarter.””

3. Amendment to Section 7.10.1 (Pipeline ROW Royalties).

The table in Section 7.10.1(a) is hereby amended and restated in its entirety as follows:

Aggregate Annual Net Revenues of Pipeline Products (excluding AMG 510)	Royalty Rate
For that portion of aggregate annual Net Revenues of Pipeline Products in the ROW less than or equal to [*] U.S. Dollars (US\$[*])	[*]%*
For that portion of aggregate annual Net Revenues of Pipeline Products in the ROW greater than [*] U.S. Dollars (US\$[*]) and less than or equal to [*] U.S. Dollars (US\$[*])	[*]%*
For that portion of aggregate annual Net Revenues of Pipeline Products in the ROW greater than [*] U.S. Dollars (US\$[*]) and less than or equal to [*] U.S. Dollars (US\$[*])	[*]%*
For that portion of aggregate annual Net Revenues of Pipeline Products in the ROW greater than [*] Dollars (US\$[*])	[*]%*
On and after January 1, 2026, the portion of aggregate annual Net Revenues of Pipeline Products attributed to Net Revenues of AMG 757 (also known as tarlatamab or IMDELLTRA®) for any Royalty Rate tier will be paid at the specified Royalty Rate for such tier reduced by []% (specifically [*]). If aggregate annual Net Revenues include Net Revenues from any Pipeline Products other than AMG 757, Net Revenues for each Pipeline Product, including AMG 757, will be distributed proportionately (based on the percentage of Net Revenues attributable to each applicable Pipeline Product) across each Royalty Rate tier. For example, if in a particular calendar year, sales of AMG 757 were \$[*] and sales of other Pipeline Products were \$[*], then the percentage applicable to AMG 757 would be [*]% and the percentage applicable to the other Pipeline Products would be [*]%. So, total royalties due for that year would be [*]. Quarterly proportions will be determined based on the cumulative year-to-date Net Revenues for each applicable Pipeline Product. Any adjustments to previous quarters resulting from changes in these cumulative proportions will be applied in the current quarter.	

4. Miscellaneous.

- a. Except as specifically amended above, the Agreement shall continue to be in full force and effect.
- b. This Amendment and its effect are subject to and shall be construed and enforced in accordance with the laws of the State of New York, U.S.A.
- c. This Amendment may be executed in counterparts with the same effect as if both Parties had signed the same document. All such counterparts will be deemed an original, will be construed together and will constitute one and the same instrument. Signature pages of this Amendment may be exchanged by facsimile or other electronic means without affecting the validity thereof.

IN WITNESS WHEREOF, the Parties have executed this Amendment as of the Fourth Amendment Effective Date.

AMGEN INC.

/s/ Robert A. Bradway

Name: Robert A. Bradway

Title: Chairman of the Board, President and
Chief Executive Officer

BEONE MEDICINES I GMBH

/s/ Michael Schoen

Name: Michael Schoen

Title: Managing Director

BEONE MEDICINES LTD.

/s/ Chan Lee

Name: Chan Lee

Title: Senior Vice President, General Counsel & Corporate Secretary

AMGEN INC.

The following is a list of subsidiaries of the Company as of December 31, 2025, omitting some subsidiaries which, considered in the aggregate, would not constitute a significant subsidiary.

SUBSIDIARY (Name under which subsidiary does business)	STATE OR OTHER JURISDICTION OF INCORPORATION OR ORGANIZATION
Amgen (Europe) GmbH	Switzerland
Amgen Canada Inc.	Canada
Amgen Fremont Inc.	Delaware
Amgen Global Finance B.V.	Netherlands
Amgen Global Technology Unlimited Company	Ireland
Amgen GmbH Germany	Germany
Amgen İlaç Ticaret Limited Şirketi	Türkiye
Amgen International Holdings Inc.	Delaware
Amgen K.K.	Japan
Amgen K-A, Inc.	Delaware
Amgen Manufacturing Limited LLC	Puerto Rico
Amgen Rare Disease Holdings Inc.	Delaware
Amgen Research (Munich) GmbH	Germany
Amgen S.A.S.	France
Amgen S.p.A.	Italy
Amgen SF, LLC	Delaware
Amgen Singapore Holding Pte. Ltd.	Singapore
Amgen Singapore Manufacturing Pte. Ltd.	Singapore
Amgen Technology (Ireland) Unlimited Company	Ireland
Amgen Technology, Limited	Bermuda
Amgen USA Inc.	Delaware
Amgen Worldwide Holdings B.V.	Netherlands
Amgen, S.A.	Spain
BioVex LLC	Delaware
ChemoCentryx, Inc.	Delaware
Five Prime Therapeutics, Inc.	Delaware
Horizon Therapeutics Finance S.à.r.l	Luxembourg
Horizon Therapeutics Holdings Limited	Ireland
Horizon Therapeutics Ireland DAC	Ireland
Horizon Therapeutics Limited	Ireland
Horizon Therapeutics Services LLC	Delaware
Horizon Therapeutics U.S. Holding LLC	Delaware
Horizon Therapeutics USA, Inc.	Delaware
HZNP Finance Limited	Bermuda
Immunex Corporation	Washington
Immunex Rhode Island Corporation	Delaware
Onyx Pharmaceuticals, Inc.	Delaware
Onyx Therapeutics, Inc.	Delaware
Pillartree Limited	Ireland
Saga Investments Coöperatief U.A.	Netherlands
TeneoBio, Inc.	Delaware
Viola Bio, Inc.	Delaware

CERTIFICATIONS

I, Robert A. Bradway, Chairman of the Board, Chief Executive Officer and President of Amgen Inc., certify that:

1. I have reviewed this Annual Report on Form 10-K of Amgen Inc.;
2. Based on my knowledge, this annual report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this annual report;
3. Based on my knowledge, the financial statements, and other financial information included in this annual report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this annual report;
4. The registrant's other certifying officer(s) and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:
 - (a) Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this annual report is being prepared;
 - (b) Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;
 - (c) Evaluated the effectiveness of the registrant's disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this annual report based on such evaluation; and
 - (d) Disclosed in this annual report any change in the registrant's internal control over financial reporting that occurred during the registrant's most recent fiscal quarter (the registrant's fourth fiscal quarter in the case of an annual report) that has materially affected, or is reasonably likely to materially affect, the registrant's internal control over financial reporting; and
5. The registrant's other certifying officer(s) and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrant's auditors and the audit committee of the registrant's board of directors (or persons performing the equivalent functions):
 - (a) All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrant's ability to record, process, summarize and report financial information; and
 - (b) Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrant's internal control over financial reporting.

February 13, 2026

/s/ ROBERT A. BRADWAY

Robert A. Bradway
Chairman of the Board,
Chief Executive Officer and President

CERTIFICATIONS

I, Peter H. Griffith, Executive Vice President and Chief Financial Officer of Amgen Inc., certify that:

1. I have reviewed this Annual Report on Form 10-K of Amgen Inc.;
2. Based on my knowledge, this annual report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this annual report;
3. Based on my knowledge, the financial statements, and other financial information included in this annual report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this annual report;
4. The registrant's other certifying officer(s) and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:
 - (a) Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this annual report is being prepared;
 - (b) Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;
 - (c) Evaluated the effectiveness of the registrant's disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this annual report based on such evaluation; and
 - (d) Disclosed in this annual report any change in the registrant's internal control over financial reporting that occurred during the registrant's most recent fiscal quarter (the registrant's fourth fiscal quarter in the case of an annual report) that has materially affected, or is reasonably likely to materially affect, the registrant's internal control over financial reporting; and
5. The registrant's other certifying officer(s) and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrant's auditors and the audit committee of the registrant's board of directors (or persons performing the equivalent functions):
 - (a) All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrant's ability to record, process, summarize and report financial information; and
 - (b) Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrant's internal control over financial reporting.

February 13, 2026

/s/ PETER H. GRIFFITH

Peter H. Griffith

Executive Vice President and Chief Financial Officer

Certification of Chief Executive Officer

Pursuant to 18 U.S.C. § 1350, as created by Section 906 of the Sarbanes-Oxley Act of 2002, the undersigned officer of Amgen Inc. (the “Company”) hereby certifies that:

- (i) the accompanying Annual Report on Form 10-K of the Company for the period ended December 31, 2025 (the “Report”) fully complies with the requirements of Section 13(a) or Section 15(d), as applicable, of the Securities Exchange Act of 1934, as amended; and
- (ii) information contained in the Report fairly presents, in all material respects, the financial condition and results of operations of the Company.

February 13, 2026

/s/ ROBERT A. BRADWAY

Robert A. Bradway
Chairman of the Board,
Chief Executive Officer and President

A signed original of this written statement required by Section 906 of the Sarbanes-Oxley Act of 2002 (“Section 906”), or other document authenticating, acknowledging, or otherwise adopting the signature that appears in typed form within the electronic version of this written statement required by Section 906, has been provided to Amgen Inc. and will be retained by Amgen Inc. and furnished to the Securities and Exchange Commission or its staff upon request.

Certification of Chief Financial Officer

Pursuant to 18 U.S.C. § 1350, as created by Section 906 of the Sarbanes-Oxley Act of 2002, the undersigned officer of Amgen Inc. (the “Company”) hereby certifies that:

- (i) the accompanying Annual Report on Form 10-K of the Company for the period ended December 31, 2025 (the “Report”) fully complies with the requirements of Section 13(a) or Section 15(d), as applicable, of the Securities Exchange Act of 1934, as amended; and
- (ii) information contained in the Report fairly presents, in all material respects, the financial condition and results of operations of the Company.

February 13, 2026

/s/ PETER H. GRIFFITH

Peter H. Griffith

Executive Vice President and Chief Financial Officer

A signed original of this written statement required by Section 906 of the Sarbanes-Oxley Act of 2002 (“Section 906”), or other document authenticating, acknowledging, or otherwise adopting the signature that appears in typed form within the electronic version of this written statement required by Section 906, has been provided to Amgen Inc. and will be retained by Amgen Inc. and furnished to the Securities and Exchange Commission or its staff upon request.